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(71) Applicant (for all designated States except US): CURAGEN CORPORATION [US/US]; 555 Long Wharf Drive, 11th Floor, New Haven, CT 06511 (US).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): SHIMKETS, Richard, A. [US/US]; 191 Leete Street, West Haven, CT 06516 (US). LEACH, Martin [GB/US]; 884 School Street, Webster, MA 01570 (US).
- (74) Agent: ELRIFI, Ivor, R.; Mintz, Levin, Cohn, Ferris, Glovsky and Popeo, P.C., One Financial Center, Boston, MA 02111 (US).
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(54) Title: NUCLEIC ACIDS INCLUDING OPEN READING FRAMES ENCODING POLYPEPTIDES; "ORFX"

(57) Abstract

The present invention provides open reading frames ORFX, encoding isolated polypeptides, as well as polynucleotides encoding ORFX and antibodies that immunospecifically bind to ORFX or any derivative, variant, mutant, or fragment of the ORFX polypeptides, polynucleotides or antibodies. The invention additionally provides methods in which the ORFX polypeptide, polynucleotide and antibody are used in detection and treatment of a broad range of pathological states, as well as to other uses.

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NOVEL POLYNUCLEOTIDES AND POLYPEPTIDES ENCODED THEREBY

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BACKGROUND OF THE INVENTION

The invention relates generally to nucleic acids and polypeptides encoded thereby, and methods of using these nucleic acids and polypeptides.

SUMMARY OF THE INVENTION

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The invention is based in part on the discovery of nucleic acids that include open reading frames encoding novel polypeptides, and on the polypeptides encoded thereby. The nucleic acids and polypeptides are collectively referred to herein as "ORFX".

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Accordingly, in one aspect, the invention provides an isolated nucleic acid molecule (SEQ ID NO:2n-1, wherein n is an integer between 1-3161), that encodes novel polypeptide, or a fragment, homolog, analog or derivative thereof. The nucleic acid can include, e.g., a nucleic acid sequence encoding a polypeptide at least 85% identical to a polypeptide comprising the amino acid sequences of SEQ ID NO:2n, wherein n is an integer between 1-3161. The nucleic acid can be, e.g., a genomic DNA fragment, or a cDNA molecule.

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Also included in the invention is a vector containing one or more of the nucleic acids described herein, and a cell containing the vectors or nucleic acids described herein.

The invention is also directed to host cells transformed with a recombinant expression vector comprising any of the nucleic acid molecules described above.

In another aspect, the invention includes a pharmaceutical composition that includes an ORFX nucleic acid and a pharmaceutically acceptable carrier or diluent.

In a further aspect, the invention includes a substantially purified ORF polypeptide, e.g., any of the ORFX polypeptides encoded by an ORFX nucleic acid, and fragments, homologs, analogs, and derivatives thereof. The invention also includes a pharmaceutical composition that includes a ORFX polypeptide and a pharmaceutically acceptable carrier or diluent.

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In a still a further aspect, the invention provides an antibody that binds specifically to an ORFX polypeptide. The antibody can be, e.g., a monoclonal or polyclonal antibody, and fragments, homologs, analogs, and derivatives thereof. The invention also includes a pharmaceutical composition including ORFX antibody and a pharmaceutically acceptable carrier or diluent. The invention is also directed to isolated antibodies that bind to an epitope on a polypeptide encoded by any of the nucleic acid molecules described above.

The invention also includes kits comprising any of the pharmaceutical compositions described above.

The invention further provides a method for producing an ORFX polypeptide by providing a cell containing a ORFX nucleic acid, e.g., a vector that includes a ORFX nucleic acid, and culturing the cell under conditions sufficient to express the ORFX polypeptide encoded by the nucleic acid. The expressed ORFX polypeptide is then recovered from the cell. Preferably, the cell produces little or no endogenous ORFX polypeptide. The cell can be, e.g., a prokaryotic cell or eukaryotic cell.

The invention is also directed to methods of identifying an ORFX polypeptide or nucleic acids in a sample by contacting the sample with a compound that specifically binds to the polypeptide or nucleic acid, and detecting complex formation, if present.

The invention further provides methods of identifying a compound that modulates the activity of a ORFX polypeptide by contacting ORFX polypeptide with a compound and determining whether the ORFX polypeptide activity is modified.

The invention is also directed to compounds that modulate ORFX polypeptide activity identified by contacting a ORFX polypeptide with the compound and determining whether the compound modifies activity of the ORFX polypeptide, binds to the ORFX polypeptide, or binds to a nucleic acid molecule encoding a ORFX polypeptide.

In a another aspect, the invention provides a method of determining the presence of or predisposition of an ORFX-associated disorder in a subject. The method includes providing a sample from the subject and measuring the amount of ORFX polypeptide in the subject sample.

The amount of ORFX polypeptide in the subject sample is then compared to the amount of ORFX polypeptide in a control sample. An alteration in the amount of ORFX polypeptide in the subject protein sample relative to the amount of ORFX polypeptide in the control protein sample indicates the subject has a tissue proliferation-associated condition. A control sample is preferably taken from a matched individual, *i.e.*, an individual of similar age, sex, or other general condition but who is not suspected of having a tissue proliferation-associated condition. Alternatively, the control sample may be taken from the subject at a time when the subject is not suspected of having a tissue proliferation-associated disorder. In some embodiments, the ORFX is detected using a ORFX antibody.

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In a further aspect, the invention provides a method of determining the presence of or predisposition of an ORFX-associated disorder in a subject. The method includes providing a nucleic acid sample, e.g., RNA or DNA, or both, from the subject and measuring the amount of the ORFX nucleic acid in the subject nucleic acid sample. The amount of ORFX nucleic acid sample in the subject nucleic acid is then compared to the amount of an ORFX nucleic acid in a control sample. An alteration in the amount of ORFX nucleic acid in the sample relative to the amount of ORFX in the control sample indicates the subject has a tissue proliferation-associated disorder.

In a still further aspect, the invention provides method of treating or preventing or delaying a ORFX-associated disorder. The method includes administering to a subject in which such treatment or prevention or delay is desired a ORFX nucleic acid, a ORFX polypeptide, or an ORFX antibody in an amount sufficient to treat, prevent, or delay a tissue proliferation-associated disorder in the subject.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described below. All publications, patent applications, patents, and other references mentioned herein are incorporated by reference in their entirety. In the case of conflict, the present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

Other features and advantages of the invention will be apparent from the following detailed description and claims.

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DETAILED DESCRIPTION OF THE INVENTION

The invention provides novel polypeptides and nucleotides encoded thereby. The polynucleotides and their encoded polypeptides can be grouped according to the functions played by their gene products. Such functions include, structural proteins, proteins from which associated with metabolic pathways fatty acid metabolism, glycolysis, intermediary metabolism, calcium metabolism, proteases, and amino acid metabolism, etc.

Included in the invention are 3161 novel nucleic acid sequences and their encoded polypeptides. The sequences are collectively referred to as "ORFX nucleic acids" or ORFX polynucleotides" and the corresponding encoded polypeptide is referred to as a "ORFX polypeptide" or ORFX protein". For example, an ORFX nucleic acid according to the invention is a nucleic acid including an ORF1 nucleic acid, and an ORF polypeptide according to the invention is a polypeptide that includes the amino acid sequence of an ORF1 polypeptide. Unless indicated otherwise, "ORFX" is meant to refer to any of the ORF1-3161 sequences disclosed herein.

Table 1 provides a summary of the ORFX nucleic acids and their encoded polypeptides are summarized in Table 1. Nucleic acid sequences and polypeptide sequences for ORFX nucleic acids according to the invention is provided in the section of the specification entitled "Disclosed Sequences of ORFX Nucleic Acid and Polypeptide Sequences."

Column 1 of Table 1, entitled "ORF #", denotes an ORF number assigned to a nucleic acid containing an open reading frame according to the invention.

Column 2 of Table 1, entitled "Internal Identification number (Nucleic Acid Sequence Identification Number, Polypeptide Sequence Identification Number), provides an internal identification number for the indicated ORF, along with sequence identification numbers (SEQ ID NOs.) corresponding to the indicated ORF. In general, for an ORFn according to the invention (wherein n is any integer from 1 to 3161), a nucleic acid corresponding to the ORF is SEQ ID NO:2n-1, and an amino acid sequence encoded by the ORF is SEQ ID NO:2n. For example, a nucleic acid sequence corresponding to an ORF1 nucleic acid is SEQ ID NO:1, and a polypeptide sequence corresponding to an ORF1 polypeptide is SEQ ID NO:2. Similarly, a

nucleic acid sequence corresponding to an ORF4 nucleic acid is SEQ ID NO:7, and a polypeptide sequence corresponding to an ORF4 polypeptide is SEQ ID NO:8; a nucleic acid sequence corresponding to an ORF198 nucleic acid sequence is SEQ ID NO:395, and a polypeptide sequence corresponding to an ORF198 polypeptide is SEQ ID NO:396. Nucleic acid sequences and polypeptide sequences for ORFX nucleic acids according to the invention are provided in the section of the specification entitled "Disclosed Sequences of ORFX Nucleic Acid and Polypeptide Sequences."

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Column 2 of Table 1, entitled "Protein Similarity", lists previously described proteins that are related to polypeptides encoded by the ORFs. Genbank identifiers for the previously described proteins are provided. These can be retrieved from http://www.ncbi.nlm.nih.gov/.

To determine similarity to previously described proteins, polypeptides encoded by ORFX DNA sequences were tested using the Framesearch Algorithm against a nonredundant version of the GenPept Database from NCBI/Genbank. DNA sequences that had a score of '90' or above (Framesearch algorithm score, Edelman et. al. GCG Genetics) to a known protein were selected. Open reading frames were extended beyond the region of the protein matched using standard DNA translation and codon tables. Novel proteins that lacked a protein match were translated against the standard genetic codons and proteins with an ORF at least 80 amino acids and containing a Methionine start are included in the Table.

Column 3 of Table 3, entitled "Protein Domains", lists previously described protein domains, designated by pfam entries, that are present in polypeptides encoded by the ORFs, Also included in column 3 are proteins in which these domains are present. The pfam entries can be retrieved from http://pfam.wustl.edu/. DNA sequences were translated in all six frames and tested using the Hmmer Algorithm against the Pfam Database (References to the algorithm and Pfam database can be found at http://pfam.wustl.edu). Translated DNA sequences that matched a protein domain entry in the Pfam database AND had a score of 7.5' were selected.

Column 4 of Table 3, entitled "Protein Classification", lists the type of classification assigned for the protein, based on its homology. Examples of proteins in the classification include the following proteins:

Amylases

Amylase is responsible for endohydrolysis of 1,4-alpha-glucosidic linkages in oligosaccharides and polysaccharides. Variations in amylase gene may be indicative of delayed maturation and of various amylase producing neoplasms and carcinomas.

Amyloid

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The serum amyloid A (SAA) proteins comprise a family of vertebrate proteins that associate predominantly with high density lipoproteins (HDL). The synthesis of certain members of the family is greatly increased in inflammation. Prolonged elevation of plasma SAA levels, as in chronic inflammation, 15 results in a pathological condition, called amyloidosis, which affects the liver, kidney and spleen and which is characterized by the highly insoluble accumulation of SAA in these tissues. Amyloid selectively inhibits insulin-stimulated glucose utilization and glycogen deposition in muscle, while not affecting adipocyte glucose metabolism. Deposition of fibrillar amyloid proteins intraneuronally, as neurofibrillary tangles, extracellularly, as plaques and in blood vessels, is characteristic of both Alzheimer's disease and aged Down's syndrome. Amyloid deposition is also associated with type II diabetes mellitus.

Angiopoeitin

Members of the angiopoeitin/fibrinogen family have been shown to stimulate the generation of new blood vessels, inhibit the generation of new blood vessels, and perform several roles in blood clotting. This generation of new blood vessels, called angiogenesis, is also an essential step in tumor growth in order for the tumor to get the blood supply it needs to expand. Variation in these genes may be predictive of any form of heart disease, numerous blood clotting disorders, stroke, hypertension and predisposition to tumor formation and metastasis. In particular, these variants may be predictive of the response to various antihypertensive drugs and chemotherapeutic and anti-tumor agents.

Apoptosis-related proteins

Active cell suicide (apoptosis) is induced by events such as growth factor withdrawal and toxins. It is controlled by regulators, which have either an inhibitory effect on programmed cell

death (anti-apoptotic) or block the protective effect of inhibitors (pro-apoptotic). Many viruses have found a way of countering defensive apoptosis by encoding their own anti-apoptosis genes preventing their target-cells from dying too soon. Variants of apoptosis related genes may be useful in formulation of anti-aging drugs.

Cadherin, Cyclin, Polymerase, Oncogenes, Histones, Kinases

Members of the cell division/cell cycle pathways such as cyclins, many transcription factors and kinases, DNA polymerases, histones, helicases and other oncogenes play a critical role in carcinogenesis where the uncontrolled proliferation of cells leads to tumor formation and eventually metastasis. Variation in these genes may be predictive of predisposition to any form of cancer, from increased risk of tumor formation to increased rate of metastasis. In particular, these variants may be predictive of the response to various chemotherapeutic and anti-tumor agents.

Colony-stimulating factor-related proteins

Granulocyte/macrophage colony-stimulating factors are cytokines that act in hematopoiesis by controlling the production, differentiation, and function of 2 related white cell populations of the blood, the granulocytes and the monocytes-macrophages.

Complement-related proteins

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Complement proteins are immune associated cytotoxic agents, acting in a chain reaction to exterminate target cells to that were opsonized (primed) with antibodies, by forming a membrane attack complex (MAC). The mechanism of killing is by opening pores in the target cell membrane. Variations in 20 complement genes or their inhibitors are associated with many autoimmune disorders. Modified serum levels of complement products cause edemas of various tissues, lupus (SLE), vasculitis, glomerulonephritis, renal failure, hemolytic anemia, thrombocytopenia, and arthritis. They interfere with mechanisms of ADCC (antibody dependent cell cytotoxicity), severely impair immune competence and reduce phagocytic ability. Variants of complement genes may also be indicative of type I diabetes mellitus, meningitis neurological disorders such as nemaline myopathy, neonatal hypotonia, muscular disorders such as congenital myopathy and other diseases.

Cytochrome

The respiratory chain is a key biochemical pathway which is essential to all aerobic cells. There are five different cytochromes involved in the chain. These are heme bound proteins which serve as electron carriers. Modifications in these genes may be predictive of ataxia areflexia, dementia and myopathic and neuropathic changes in muscles. Also, association with various types of solid tumors.

Kinesins

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Kinesins are tubulin molecular motors that function to transport organelles within cells and to move chromosomes along microtubules during cell division. Modifications of these genes may be indicative of neurological disorders such as Pick disease of the brain, tuberous sclerosis.

Cytokines, Interferon, Interleukin

Members of the cytokine families are known for their potent ability to stimulate cell growth and division even at low concentrations. Cytokines such as erythropoietin are cell-specific in their growth stimulation; erythropoietin is useful for the stimulation of the proliferation of erythroblasts. Variants in cytokines may be predictive for a wide variety of diseases, including cancer predisposition.

G-protein coupled receptors

G-protein coupled receptors (also called R7G) are an extensive group of hormones, neurotransmitters, odorants and light receptors which transduce extracellular signals by interaction with guanine nucleotide-binding (G) proteins. Alterations in genes coding for G-coupled proteins may be involved in and indicative of a vast number of physiological conditions. These include blood pressure regulation, renal dysfunctions, male infertility, dopamine associated cognitive, emotional, and endocrine functions, hypercalcemia, chondrodysplasia and osteoporosis, pseudohypoparathyroidism, growth retardation and dwarfism.

Thioesterases

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Eukaryotic thiol proteases are a family of proteolytic enzymes which contain an active site cysteine. Catalysis proceeds through a thioester intermediate and is facilitated by a nearby histidine side chain; an asparagine completes the essential catalytic triad. Variants of thioester associated genes may be predictive of neuronal disorders and mental illnesses such as Ceroid Lipoffiscinosis, Neuronal 1, Infantile, Santavuori disease and more.

The key to the molecule type is as follows:

10	Abbrev:	Title:
	amylase	amylase protein
	amylaseinhib	amylase inhibitor
	amyloid	amyloid protein
15	apoptosis	apoptosis associated protein
	apoptosisinhib	apoptosis inhibitors
	apoptosisrecep	apoptosis receptors
	ATPase_associated	ATPase associated protein
	biotindep	biotin dependent enzyme/protein
20	cadherin	cadherin protein
	calcium_channel	calcium channel protein
	carboxylase	carboxylase protein
	cathepsin	cathepsin/carboxypeptidases
	cathepsininhib	cathepsin/carboxypeptidase inhibitor
25	chloride_channel	chloride channel protein
	collagen	collagen
	complement	complement protein
	complementrecept	complement receptor protein
	complementinhib	complement inhibitor
30	csf	colony stimulating factor
	csfrecept	colony stimulating factor receptor
	èyclin	cyclin protein
	cyto450	cytochrome p450 protein
	cytochrome	cytochrome related protein
35	deaminase	deaminase
	dehydrogenase	dehydrogenase
	desaturase	desaturase
	dna_rna_bind	DNA/RNA binding protein/factor
	dna_rna_inhib	DNA/RNA binding protein/factor inhibitor
40	dynein	dynein

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	elastase	elastase
	elastaseinhib	elastase inhibitor
	eph	EPH family of tyrosine kinases
	esterase	esterase
5	esteraseinhib	esterase inhibitor
	fgf	fibroblast growth factor
	fgfreceptor	fibroblast growth factor receptor
	gaba	GABA receptor
	glucoamylase	glucoamylase
10	glucoronidase	glucoronidase
	glycoprotein	glycoprotein
	Guanylyl	guanylylate cyclase
	helicase	helicase
	histone	histone
15	НОМ	homologous
	homeobox	homeobox protein
	hydrolase	hydrolase
	hydroxysteroid	hydroxysteroid associated protein
	hypoxanthine	hypoxanthine associated protein
20	immunoglob	immunoglobulin
	immunoglobrecept	immunoglobulin receptor
	interferon	interferon
	interleukin	interleukin
	interleukinrecept	interleukin receptor
25	isomerase	isomerase
	isomeraseinhibitor	isomerase inhibitor
	isomerasereceptor	isomerase receptor
	kinase	kinase
	kinaseinhibitor	kinase inhibitor
30	kinasereceptor	kinase receptor
	kinesin	kinesin
	laminin	laminin associated protein
	lipase	lipase
	metallothionein	metallothionein
35	MHC	major histocompatability complex
	misc_channel	miscellaneous channel
	ngf	nerve growth factor
	nuci_recpt	nuclear receptor
	nuclease	nuclease
40	oncogene	oncogene associated protein
	oxidase	oxidase
	oxygenase	oxygenase
	peptidase	peptidase
	peroxidase	peroxidase
45	phosphatase	phosphatase
	phosphataseinhib	phosphatase inhibitor

	phosphorylase PIR	phosphorylase PIR DATABASE (release 56, 29-OCT- 1998)
	polymerase	polymerase
5	potassium_channel	potassium channel protein
	prostaglandin	prostaglandin
	protease	protease
	proteaseinhib	protease inhibitor
	reductase	reductase
10	ribosomalprot	ribosomal associated protein
	RTR	EMBLDATABASE translated entries not to
		be incorporated into SWISS-PROT (20-
		JUL-1998)
	SIM	similar
15	SPTR	EMBL DATABASE translated entries to be
		incorporated into SWISS-PROT (20-JUL-
		1998)
	struct	structural associated protein
	sulfotransferase	sulfotransferase
20	SWP	SWISS-PROT DATABASE (release 18-
	GIV TO I	OCT-1998)
	SWPN	SWISS-PROT Update (release 11-NOV-98)
	synthase	synthase
25	tgf	transforming growth factor
25	tgfreceptor	transforming growth factor receptor thioesterase
	thioesterase thiolase	thiolase
	tm7	seven transmembrane domain G-protein
	un/	coupled receptor
30	tnf	necrosis factor receptor
30	traffic	tumor necrosis factor
	tnfreceptor	tumor trafficking associated protein
	TRN	EMBL DATABASE translated entries
		update (20-JUL-1998)
35	transcriptfactor	transcription factor
33	transferase	transferase
	transport	transport protein
	tubulin	tubulin
	ubiquitin	ubiquitin
40	unclassified	Protein not categorized into one of the
· -		aforementioned protein families
	water channel	water channel protein
		•

Column 5 of Table 1, entitled, "Cells or Tissues in Which Gene is Expressed", denotes tissues, represented by five digit numbers, in which RNA homologous to the ORF nucleic acid sequences is present. Tissues or cells corresponding to the numbers are provided in Table 2.

ORFX nucleic acids, and their encoded polypeptides, according to the invention are useful in a variety of applications and contexts. For example, various ORFX nucleic acids and polypeptides according to the invention are useful, *inter alia*, as novel members of the protein families indicated in Table 1, and/or according to the presence of domains and sequence relatedness to previously described proteins as summarized in Table 1.

ORFX nucleic acids and polypeptides according to the invention can also be used to identify cell types listed in Table 1 for an indicated ORFX according to the invention.

Additional utilities for ORFX nucleic acids and polypeptides according to the invention are disclosed herein.

ORFX Nucleic Acids

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The novel nucleic acids of the invention include those that encode an ORFX or ORFX-like protein, or biologically active portions thereof. The nucleic acids include nucleic acids encoding polypeptides that include the amino acid sequence of one or more of SEQ ID NO:2n, wherein n = 1 to 3161. The encoded polypeptides can thus include, e.g., the amino acid sequences of SEQ ID NO: 2, 4, 6, 8, 10, ..., 6310, 6312, 6314, 6316, 6318, 6320, and/or 6322.

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In some embodiments, a nucleic acid encoding a polypeptide having the amino acid sequence of one or more of SEQ ID NO:2n (wherein n = 1 to 3161) includes the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a fragment thereof. Additionally, the invention includes mutant or variant nucleic acids of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a fragment thereof, any of whose bases may be changed from the disclosed sequence while still encoding a protein that maintains its ORFX -like activities and physiological functions. The invention further includes the complement of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), including fragments, derivatives,

analogs and homolog thereof. The invention additionally includes nucleic acids or nucleic acid fragments, or complements thereto, whose structures include chemical modifications.

Also included are nucleic acid fragments sufficient for use as hybridization probes to identify ORFX-encoding nucleic acids (e.g., ORFX mRNA) and fragments for use as polymerase chain reaction (PCR) primers for the amplification or mutation of ORFX nucleic acid molecules. As used herein, the term "nucleic acid molecule" is intended to include DNA molecules (e.g., cDNA or genomic DNA), RNA molecules (e.g., mRNA), analogs of the DNA or RNA generated using nucleotide analogs, and derivatives, fragments and homologs thereof. The nucleic acid molecule can be single-stranded or double-stranded, but preferably is double-stranded DNA.

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"Probes" refer to nucleic acid sequences of variable length, preferably between at least about 10 nucleotides (nt), 100 nt, or as many as about, e.g., 6,000 nt, depending on use. Probes are used in the detection of identical, similar, or complementary nucleic acid sequences. Longer length probes are usually obtained from a natural or recombinant source, are highly specific and much slower to hybridize than oligomers. Probes may be single- or double-stranded and designed to have specificity in PCR, membrane-based hybridization technologies, or ELISA-like technologies.

An "isolated" nucleic acid molecule is one that is separated from other nucleic acid molecules that are present in the natural source of the nucleic acid. Examples of isolated nucleic acid molecules include, but are not limited to, recombinant DNA molecules contained in a vector, recombinant DNA molecules maintained in a heterologous host cell, partially or substantially purified nucleic acid molecules, and synthetic DNA or RNA molecules. Preferably, an "isolated" nucleic acid is free of sequences which naturally flank the nucleic acid (*i.e.*, sequences located at the 5' and 3' ends of the nucleic acid) in the genomic DNA of the organism from which the nucleic acid is derived. For example, in various embodiments, the isolated ORFX nucleic acid molecule can contain less than about 50 kb, 25 kb, 5 kb, 4 kb, 3 kb, 2 kb, 1 kb, 0.5 kb or 0.1 kb of nucleotide sequences which naturally flank the nucleic acid molecule in genomic DNA of the cell from which the nucleic acid is derived. Moreover, an "isolated" nucleic acid molecule, such as a cDNA molecule, can be substantially free of other cellular material or culture medium when produced by recombinant techniques, or of chemical precursors or other chemicals when chemically synthesized.

A nucleic acid molecule of the present invention, e.g., a nucleic acid molecule having the nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a complement of any of this nucleotide sequence, can be isolated using standard molecular biology techniques and the sequence information provided herein. Using all or a portion of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161) as a hybridization probe, ORFX nucleic acid sequences can be isolated using standard hybridization and cloning techniques (e.g., as described in Sambrook et al., eds., Molecular Cloning: A Laboratory Manual 2^{nd} Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989; and Ausubel, et al., eds., Current Protocols in Molecular Biology, John Wiley & Sons, New York, NY, 1993.)

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A nucleic acid of the invention can be amplified using cDNA, mRNA or alternatively, genomic DNA, as a template and appropriate oligonucleotide primers according to standard PCR amplification techniques. The nucleic acid so amplified can be cloned into an appropriate vector and characterized by DNA sequence analysis. Furthermore, oligonucleotides corresponding to ORFX nucleotide sequences can be prepared by standard synthetic techniques, e.g., using an automated DNA synthesizer.

As used herein, the term "oligonucleotide" refers to a series of linked nucleotide residues, which oligonucleotide has a sufficient number of nucleotide bases to be used in a PCR reaction. A short oligonucleotide sequence may be based on, or designed from, a genomic or cDNA sequence and is used to amplify, confirm, or reveal the presence of an identical, similar or complementary DNA or RNA in a particular cell or tissue. Oligonucleotides comprise portions of a nucleic acid sequence having about 10 nt, 50 nt, or 100 nt in length, preferably about 15 nt to 30 nt in length. In one embodiment, an oligonucleotide comprising a nucleic acid molecule less than 100 nt in length would further comprise at lease 6 contiguous nucleotides of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a complement thereof. Oligonucleotides may be chemically synthesized and may be used as probes.

In another embodiment, an isolated nucleic acid molecule of the invention comprises a nucleic acid molecule that is a complement of the nucleotide sequence shown in any of SEQ ID NO:2n-1 (wherein n=1 to 3161). In another embodiment, an isolated nucleic acid molecule of the invention comprises a nucleic acid molecule that is a complement of the nucleotide sequence shown in any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a portion of this nucleotide sequence. A nucleic acid molecule that is complementary to the nucleotide sequence shown in

is one that is sufficiently complementary to the nucleotide sequence shown in of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161) that it can hydrogen bond with little or no mismatches to the nucleotide sequence shown in of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161),, thereby forming a stable duplex.

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As used herein, the term "complementary" refers to Watson-Crick or Hoogsteen base pairing between nucleotides units of a nucleic acid molecule, and the term "binding" means the physical or chemical interaction between two polypeptides or compounds or associated polypeptides or compounds or combinations thereof. Binding includes ionic, non-ionic, Von der Waals, hydrophobic interactions, etc. A physical interaction can be either direct or indirect. Indirect interactions may be through or due to the effects of another polypeptide or compound. Direct binding refers to interactions that do not take place through, or due to, the effect of another polypeptide or compound, but instead are without other substantial chemical intermediates.

Moreover, the nucleic acid molecule of the invention can comprise only a portion of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), e.g., a fragment that can be used as a probe or primer, or a fragment encoding a biologically active portion of ORFX. Fragments provided herein are defined as sequences of at least 6 (contiguous) nucleic acids or at least 4 (contiguous) amino acids, a length sufficient to allow for specific hybridization in the case of nucleic acids or for specific recognition of an epitope in the case of amino acids, respectively, and are at most some portion less than a full length sequence. Fragments may be derived from any contiguous portion of a nucleic acid or amino acid sequence of choice. Derivatives are nucleic acid sequences or amino acid sequences formed from the native compounds either directly or by modification or partial substitution. Analogs are nucleic acid sequences or amino acid sequences that have a structure similar to, but not identical to, the native compound but differs from it in respect to certain components or side chains. Analogs may be synthetic or from a different evolutionary origin and may have a similar or opposite metabolic activity compared to wild type.

Derivatives and analogs may be full length or other than full length, if the derivative or analog contains a modified nucleic acid or amino acid, as described below. Derivatives or analogs of the nucleic acids or proteins of the invention include, but are not limited to, molecules comprising regions that are substantially homologous to the nucleic acids or proteins of the

invention, in various embodiments, by at least about 70%, 80%, 85%, 90%, 95%, 98%, or even 99% identity (with a preferred identity of 80-99%) over a nucleic acid or amino acid sequence of identical size or when compared to an aligned sequence in which the alignment is done by a computer homology program known in the art, or whose encoding nucleic acid is capable of hybridizing to the complement of a sequence encoding the aforementioned proteins under stringent, moderately stringent, or low stringent conditions. See *e.g.* Ausubel, *et al.*, Current Protocols in Molecular Biology, John Wiley & Sons, New York, NY, 1993, and below. An exemplary program is the Gap program (Wisconsin Sequence Analysis Package, Version 8 for UNIX, Genetics Computer Group, University Research Park, Madison, WI) using the default settings, which uses the algorithm of Smith and Waterman (Adv. Appl. Math., 1981, 2: 482-489, which is incorporated herein by reference in its entirety).

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A "homologous nucleic acid sequence" or "homologous amino acid sequence," or variations thereof, refer to sequences characterized by a homology at the nucleotide level or amino acid level as discussed above. Homologous nucleotide sequences encode those sequences coding for isoforms of ORFX polypeptide. Isoforms can be expressed in different tissues of the same organism as a result of, for example, alternative splicing of RNA. Alternatively, isoforms can be encoded by different genes. In the present invention, homologous nucleotide sequences include nucleotide sequences encoding for a ORFX polypeptide of species other than humans, including, but not limited to, mammals, and thus can include, e.g., mouse, rat, rabbit, dog, cat cow, horse, and other organisms. Homologous nucleotide sequences also include, but are not limited to, naturally occurring allelic variations and mutations of the nucleotide sequences set forth herein. A homologous nucleotide sequence does not, however, include the nucleotide sequence encoding human ORFX protein. Homologous nucleic acid sequences include those nucleic acid sequences that encode conservative amino acid substitutions (see below) in any of SEQ ID NO:2n (wherein n = 1 to 3161) as well as a polypeptide having ORFX activity. Biological activities of the ORFX proteins are described below. A homologous amino acid sequence does not encode the amino acid sequence of a human ORFX polypeptide.

The nucleotide sequence determined from the cloning of the human ORFX gene allows for the generation of probes and primers designed for use in identifying the cell types disclosed and/or cloning ORFX homologues in other cell types, e.g., from other tissues, as well as ORFX homologues from other mammals. The probe/primer typically comprises a substantially purified

oligonucleotide. The oligonucleotide typically comprises a region of nucleotide sequence that hybridizes under stringent conditions to at least about 12, 25, 50, 100, 150, 200, 250, 300, 350 or 400 or more consecutive sense strand nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161); or an anti-sense strand nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161); or of a naturally occurring mutant of SEQ ID NO:2n-1 (wherein n = 1 to 3161).

Probes based on the human ORFX nucleotide sequence can be used to detect transcripts or genomic sequences encoding the same or homologous proteins. In various embodiments, the probe further comprises a label group attached thereto, e.g., the label group can be a radioisotope, a fluorescent compound, an enzyme, or an enzyme co-factor. Such probes can be used as a part of a diagnostic test kit for identifying cells or tissue which misexpress a ORFX protein, such as by measuring a level of a ORFX-encoding nucleic acid in a sample of cells from a subject e.g., detecting ORFX mRNA levels or determining whether a genomic ORFX gene has been mutated or deleted.

"A polypeptide having a biologically active portion of ORFX" refers to polypeptides exhibiting activity similar, but not necessarily identical to, an activity of a polypeptide of the present invention, including mature forms, as measured in a particular biological assay, with or without dose dependency. A nucleic acid fragment encoding a "biologically active portion of ORFX" can be prepared by isolating a portion of SEQ ID NO:2n-1 (wherein n=1 to 3161), that encodes a polypeptide having a ORFX biological activity (biological activities of the ORFX proteins are summarized in Table 1), expressing the encoded portion of ORFX protein (e.g., by recombinant expression *in vitro*) and assessing the activity of the encoded portion of ORFX. For example, a nucleic acid fragment encoding a biologically active portion of ORFX can optionally include a domain as shown in Table 1, column 4.

ORFX variants

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The invention further encompasses nucleic acid molecules that differ from the disclosed ORFX nucleotide sequences due to degeneracy of the genetic code. These nucleic acids thus encode the same ORFX protein as that encoded by the nucleotide sequence shown in SEQ ID NO:2n-1 (wherein n=1 to 3161). In another embodiment, an isolated nucleic acid molecule of the invention has a nucleotide sequence encoding a protein having an amino acid sequence shown in any of SEQ ID NO:2n (wherein n=1 to 3161).

In addition to the human ORFX nucleotide sequence shown in any of SEQ ID NO:2*n*-1 (wherein n = 1 to 3161), it will be appreciated by those skilled in the art that DNA sequence polymorphisms that lead to changes in the amino acid sequences of ORFX may exist within a population (*e.g.*, the human population). Such genetic polymorphism in the ORFX gene may exist among individuals within a population due to natural allelic variation. As used herein, the terms "gene" and "recombinant gene" refer to nucleic acid molecules comprising an open reading frame encoding a ORFX protein, preferably a mammalian ORFX protein. Such natural allelic variations can typically result in 1-5% variance in the nucleotide sequence of the ORFX gene. Any and all such nucleotide variations and resulting amino acid polymorphisms in ORFX that are the result of natural allelic variation and that do not alter the functional activity of ORFX are intended to be within the scope of the invention.

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Moreover, nucleic acid molecules encoding ORFX proteins from other species, and thus that have a nucleotide sequence that differs from the human sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), are intended to be within the scope of the invention. Nucleic acid molecules corresponding to natural allelic variants and homologues of the ORFX cDNAs of the invention can be isolated based on their homology to the human ORFX nucleic acids disclosed herein using the human cDNAs, or a portion thereof, as a hybridization probe according to standard hybridization techniques under stringent hybridization conditions.

In another embodiment, an isolated nucleic acid molecule of the invention is at least 6 nucleotides in length and hybridizes under stringent conditions to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2*n*-1 (wherein n = 1 to 3161). In another embodiment, the nucleic acid is at least 10, 25, 50, 100, 250, 500 or 750 nucleotides in length. In another embodiment, an isolated nucleic acid molecule of the invention hybridizes to the coding region. As used herein, the term "hybridizes under stringent conditions" is intended to describe conditions for hybridization and washing under which nucleotide sequences at least 60% homologous to each other typically remain hybridized to each other.

Homologs (i.e., nucleic acids encoding ORFX proteins derived from species other than human) or other related sequences (e.g., paralogs) can be obtained by low, moderate or high stringency hybridization with all or a portion of the particular human sequence as a probe using methods well known in the art for nucleic acid hybridization and cloning.

As used herein, the phrase "stringent hybridization conditions" refers to conditions under which a probe, primer or oligonucleotide will hybridize to its target sequence, but to no other sequences. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures than shorter sequences. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH. The Tm is the temperature (under defined ionic strength, pH and nucleic acid concentration) at which 50% of the probes complementary to the target sequence hybridize to the target sequence at equilibrium. Since the target sequences are generally present at excess, at Tm, 50% of the probes are occupied at equilibrium. Typically, stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes, primers or oligonucleotides (e.g., 10 nt to 50 nt) and at least about 60°C for longer probes, primers and oligonucleotides. Stringent conditions may also be achieved with the addition of destabilizing agents, such as formamide.

Stringent conditions are known to those skilled in the art and can be found in CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. Preferably, the conditions are such that sequences at least about 65%, 70%, 75%, 85%, 90%, 95%, 98%, or 99% homologous to each other typically remain hybridized to each other. A non-limiting example of stringent hybridization conditions is hybridization in a high salt buffer comprising 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500 mg/ml denatured salmon sperm DNA at 65°C. This hybridization is followed by one or more washes in 0.2X SSC, 0.01% BSA at 50°C. An isolated nucleic acid molecule of the invention that hybridizes under stringent conditions to the sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161) corresponds to a naturally occurring nucleic acid molecule. As used herein, a "naturally-occurring" nucleic acid molecule refers to an RNA or DNA molecule having a nucleotide sequence that occurs in nature (e.g., encodes a natural protein).

In a second embodiment, a nucleic acid sequence that is hybridizable to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or fragments, analogs or derivatives thereof, under conditions of moderate stringency is provided. A non-limiting example of moderate stringency hybridization conditions are

hybridization in 6X SSC, 5X Denhardt's solution, 0.5% SDS and 100 mg/ml denatured salmon sperm DNA at 55°C, followed by one or more washes in 1X SSC, 0.1% SDS at 37°C. Other conditions of moderate stringency that may be used are well known in the art. See, e.g., Ausubel et al. (eds.), 1993, CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, NY, and Kriegler, 1990, GENE TRANSFER AND EXPRESSION, A LABORATORY MANUAL, Stockton Press, NY.

In a third embodiment, a nucleic acid that is hybridizable to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2*n*-1 (wherein *n* = 1 to 3161), or fragments, analogs or derivatives thereof, under conditions of low stringency, is provided. A non-limiting example of low stringency hybridization conditions are hybridization in 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.2% BSA, 100 mg/ml denatured salmon sperm DNA, 10% (wt/vol) dextran sulfate at 40°C, followed by one or more washes in 2X SSC, 25 mM Tris-HCl (pH 7.4), 5 mM EDTA, and 0.1% SDS at 50°C. Other conditions of low stringency that may be used are well known in the art (*e.g.*, as employed for cross-species hybridizations). See, *e.g.*, Ausubel *et al.* (eds.), 1993, CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, NY, and Kriegler, 1990, GENE TRANSFER AND EXPRESSION, A LABORATORY MANUAL, Stockton Press, NY; Shilo and Weinberg, 1981, *Proc Natl Acad Sci USA* 78: 6789-6792.

Conservative mutations

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In addition to naturally-occurring allelic variants of the ORFX sequence that may exist in the population, the skilled artisan will further appreciate that changes can be introduced by mutation into the nucleotide sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), thereby leading to changes in the amino acid sequence of the encoded ORFX protein, without altering the functional ability of the ORFX protein. For example, nucleotide substitutions leading to amino acid substitutions at "non-essential" amino acid residues can be made in the sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161). A "non-essential" amino acid residue is a residue that can be altered from the wild-type sequence of ORFX without altering the biological activity, whereas an "essential" amino acid residue is required for biological activity. For example, amino acid residues that are conserved among the ORFX proteins of the present invention, are predicted to be particularly unamenable to alteration.

Amino acid residues that are conserved among members of an ORFX family members are predicted to be less amenable to alteration. For example, an ORFX protein according to the present invention can contain at least one domain (e.g., as shown in Table 1) that is a typically conserved region in an ORFX family member. As such, these conserved domains are not likely to be amenable to mutation. Other amino acid residues, however, (e.g., those that are not conserved or only semi-conserved among members of the ORFX family) may not be as essential for activity and thus are more likely to be amenable to alteration.

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Another aspect of the invention pertains to nucleic acid molecules encoding ORFX proteins that contain changes in amino acid residues that are not essential for activity. Such ORFX proteins differ in amino acid sequence from any of any of SEQ ID NO:2n (wherein n = 1 to 3161), yet retain biological activity. In one embodiment, the isolated nucleic acid molecule comprises a nucleotide sequence encoding a protein, wherein the protein comprises an amino acid sequence at least about 75% homologous to the amino acid sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161). Preferably, the protein encoded by the nucleic acid is at least about 80% homologous to any of SEQ ID NO:2n (wherein n = 1 to 3161), more preferably at least about 90%, 95%, 98%, and most preferably at least about 99% homologous to SEQ ID NO:2.

An isolated nucleic acid molecule encoding a ORFX protein homologous to the protein of any of SEQ ID NO:2n (wherein n = 1 to 3161) can be created by introducing one or more nucleotide substitutions, additions or deletions into the corresponding nucleotide sequence, *i.e.* SEQ ID NO:2n-1 for the corresponding n, such that one or more amino acid substitutions, additions or deletions are introduced into the encoded protein.

Mutations can be introduced into SEQ ID NO:2*n*-1 (wherein n = 1 to 3161) by standard techniques, such as site-directed mutagenesis and PCR-mediated mutagenesis. Preferably, conservative amino acid substitutions are made at one or more predicted non-essential amino acid residues. A "conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a similar side chain. Families of amino acid residues having similar side chains have been defined in the art. These families include amino acids with basic side chains (*e.g.*, lysine, arginine, histidine), acidic side chains (*e.g.*, aspartic acid, glutamic acid), uncharged polar side chains (*e.g.*, glycine, asparagine, glutamine, serine, threonine, tyrosine, cysteine), nonpolar side chains (*e.g.*, alanine, valine, leucine, isoleucine, proline,

phenylalanine, methionine, tryptophan), beta-branched side chains (e.g., threonine, valine, isoleucine) and aromatic side chains (e.g., tyrosine, phenylalanine, tryptophan, histidine). Thus, a predicted nonessential amino acid residue in ORFX is replaced with another amino acid residue from the same side chain family. Alternatively, in another embodiment, mutations can be introduced randomly along all or part of a ORFX coding sequence, such as by saturation mutagenesis, and the resultant mutants can be screened for ORFX biological activity to identify mutants that retain activity. Following mutagenesis of SEQ ID NO:2n-1 (wherein n = 1 to 3161), the encoded protein can be expressed by any recombinant technology known in the art and the activity of the protein can be determined.

In one embodiment, a mutant ORFX protein can be assayed for (1) the ability to form protein:protein interactions with other ORFX proteins, other cell-surface proteins, or biologically active portions thereof, (2) complex formation between a mutant ORFX protein and a ORFX receptor; (3) the ability of a mutant ORFX protein to bind to an intracellular target protein or biologically active portion thereof; (e.g., avidin proteins); (4) the ability to bind BRA protein; or (5) the ability to specifically bind an anti-ORFX protein antibody.

Antisense

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Another aspect of the invention pertains to isolated antisense nucleic acid molecules that are hybridizable to or complementary to the nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO:2n-1 (wherein n=1 to 3161), or fragments, analogs or derivatives thereof. An "antisense" nucleic acid comprises a nucleotide sequence that is complementary to a "sense" nucleic acid encoding a protein, e.g., complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. In specific aspects, antisense nucleic acid molecules are provided that comprise a sequence complementary to at least about 10, 25, 50, 100, 250 or 500 nucleotides or an entire ORFX coding strand, or to only a portion thereof. Nucleic acid molecules encoding fragments, homologs, derivatives and analogs of a ORFX protein of any of SEQ ID NO:2n (wherein n=1 to 3161) or antisense nucleic acids complementary to a ORFX nucleic acid sequence of SEQ ID NO:2n-1 (wherein n=1 to 3161) are additionally provided.

In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence encoding ORFX. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid

residues (e.g., the protein coding region of a human ORFX that corresponds to any of SEQ ID NO:2n (wherein n = 1 to 3161)). In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence encoding ORFX. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (i.e., also referred to as 5' and 3' untranslated regions).

Given the coding strand sequences encoding ORFX disclosed herein (e.g., SEQ ID NO:2n-1 (wherein n = 1 to 3161)), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of ORFX mRNA, but more preferably is an oligonucleotide that is antisense to only a portion of the coding or noncoding region of ORFX mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of ORFX mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis or enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used.

Examples of modified nucleotides that can be used to generate the antisense nucleic acid include: 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil,

3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (*i.e.*, RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target nucleic acid of interest, described further in the following subsection).

The antisense nucleic acid molecules of the invention are typically administered to a subject or generated in situ such that they hybridize with or bind to cellular mRNA and/or genomic DNA encoding a ORFX protein to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule that binds to DNA duplexes, through specific interactions in the major groove of the double helix. An example of a route of administration of antisense nucleic acid molecules of the invention includes direct injection at a tissue site. Alternatively, antisense nucleic acid molecules can be modified to target selected cells and then administered systemically. For example, for systemic administration, antisense molecules can be modified such that they specifically bind to receptors or antigens expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecules to peptides or antibodies that bind to cell surface receptors or antigens. The antisense nucleic acid molecules can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong pol II or pol III promoter are preferred.

In yet another embodiment, the antisense nucleic acid molecule of the invention is an α-anomeric nucleic acid molecule. An α-anomeric nucleic acid molecule forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual β-units, the strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids Res 15: 6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res 15: 6131-6148) or a chimeric RNA -DNA analogue (Inoue et al. (1987) FEBS Lett 215: 327-330).

Ribozymes and PNA moieties

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Such modifications include, by way of nonlimiting example, modified bases, and nucleic acids whose sugar phosphate backbones are modified or derivatized. These modifications are

carried out at least in part to enhance the chemical stability of the modified nucleic acid, such that they may be used, for example, as antisense binding nucleic acids in therapeutic applications in a subject.

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In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity that are capable of cleaving a single-stranded nucleic acid, such as an mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) Nature 334:585-591)) can be used to catalytically cleave ORFX mRNA transcripts to thereby inhibit translation of ORFX mRNA. A ribozyme having specificity for a ORFX-encoding nucleic acid can be designed based upon the nucleotide sequence of a ORFX DNA disclosed herein (i.e., SEQ ID NO:2n-1 (wherein n = 1 to 3161)). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is complementary to the nucleotide sequence to be cleaved in a ORFX-encoding mRNA. See, e.g., Cech et al. U.S. Pat. No. 4,987,071; and Cech et al. U.S. Pat. No. 5,116,742. Alternatively, ORFX mRNA can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, e.g., Bartel et al., (1993) Science 261:1411-1418.

Alternatively, ORFX gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region of the ORFX (e.g., the ORFX promoter and/or enhancers) to form triple helical structures that prevent transcription of the ORFX gene in target cells. See generally, Helene. (1991) Anticancer Drug Des. 6: 569-84; Helene. et al. (1992) Ann. N.Y. Acad. Sci. 660:27-36; and Maher (1992) Bioassays 14: 807-15.

In various embodiments, the nucleic acids of ORFX can be modified at the base moiety, sugar moiety or phosphate backbone to improve, e.g., the stability, hybridization, or solubility of the molecule. For example, the deoxyribose phosphate backbone of the nucleic acids can be modified to generate peptide nucleic acids (see Hyrup et al. (1996) Bioorg Med Chem 4: 5-23). As used herein, the terms "peptide nucleic acids" or "PNAs" refer to nucleic acid mimics, e.g., DNA mimics, in which the deoxyribose phosphate backbone is replaced by a pseudopeptide backbone and only the four natural nucleobases are retained. The neutral backbone of PNAs has been shown to allow for specific hybridization to DNA and RNA under conditions of low ionic strength. The synthesis of PNA oligomers can be performed using standard solid phase peptide

synthesis protocols as described in Hyrup et al. (1996) above; Perry-O'Keefe et al. (1996) PNAS 93: 14670-675.

PNAs of ORFX can be used in therapeutic and diagnostic applications. For example, PNAs can be used as antisense or antigene agents for sequence-specific modulation of gene expression by, e.g., inducing transcription or translation arrest or inhibiting replication. PNAs of ORFX can also be used, e.g., in the analysis of single base pair mutations in a gene by, e.g., PNA directed PCR clamping; as artificial restriction enzymes when used in combination with other enzymes, e.g., S1 nucleases (Hyrup B. (1996) above); or as probes or primers for DNA sequence and hybridization (Hyrup et al. (1996), above; Perry-O'Keefe (1996), above).

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In another embodiment, PNAs of ORFX can be modified, e.g., to enhance their stability or cellular uptake, by attaching lipophilic or other helper groups to PNA, by the formation of PNA-DNA chimeras, or by the use of liposomes or other techniques of drug delivery known in the art. For example, PNA-DNA chimeras of ORFX can be generated that may combine the advantageous properties of PNA and DNA. Such chimeras allow DNA recognition enzymes, e.g., RNase H and DNA polymerases, to interact with the DNA portion while the PNA portion would provide high binding affinity and specificity. PNA-DNA chimeras can be linked using linkers of appropriate lengths selected in terms of base stacking, number of bonds between the nucleobases, and orientation (Hyrup (1996) above). The synthesis of PNA-DNA chimeras can be performed as described in Hyrup (1996) above and Finn et al. (1996) Nucl Acids Res 24: 3357-63. For example, a DNA chain can be synthesized on a solid support using standard phosphoramidite coupling chemistry, and modified nucleoside analogs, e.g., 5'-(4-methoxytrityl)amino-5'-deoxy-thymidine phosphoramidite, can be used between the PNA and the 5' end of DNA (Mag et al. (1989) Nucl Acid Res 17: 5973-88). PNA monomers are then coupled in a stepwise manner to produce a chimeric molecule with a 5' PNA segment and a 3' DNA segment (Finn et al. (1996) above). Alternatively, chimeric molecules can be synthesized with a 5' DNA segment and a 3' PNA segment. See, Petersen et al. (1975) Bioorg Med Chem Lett 5: 1119-11124.

In other embodiments, the oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. W088/09810) or

the blood-brain barrier (see, e.g., PCT Publication No. W089/10134). In addition, oligonucleotides can be modified with hybridization triggered cleavage agents (See, e.g., Krol et al., 1988, BioTechniques 6:958-976) or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5: 539-549). To this end, the oligonucleotide may be conjugated to another molecule, e.g., a peptide, a hybridization triggered cross-linking agent, a transport agent, a hybridization-triggered cleavage agent, etc.

ORFX polypeptides

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The novel protein of the invention includes the ORFX-like protein whose sequence is provided in any of SEQ ID NO:2n (wherein n = 1 to 3161). The invention also includes a mutant or variant protein any of whose residues may be changed from the corresponding residue shown in FIG. 1 while still encoding a protein that maintains its ORFX-like activities and physiological functions, or a functional fragment thereof. For example, the invention includes the polypeptides encoded by the variant ORFX nucleic acids described above. In the mutant or variant protein, up to 20% or more of the residues may be so changed.

In general, an ORFX -like variant that preserves ORFX-like function includes any variant in which residues at a particular position in the sequence have been substituted by other amino acids, and further include the possibility of inserting an additional residue or residues between two residues of the parent protein as well as the possibility of deleting one or more residues from the parent sequence. Any amino acid substitution, insertion, or deletion is encompassed by the invention. In favorable circumstances, the substitution is a conservative substitution as defined above. Furthermore, without limiting the scope of the invention, positions of any of SEQ ID NO:2n (wherein n=1 to 3161) may be substitute such that a mutant or variant protein may include one or more substitutions

The invention also includes isolated ORFX proteins, and biologically active portions thereof, or derivatives, fragments, analogs or homologs thereof. Also provided are polypeptide fragments suitable for use as immunogens to raise anti-ORFX antibodies. In one embodiment, native ORFX proteins can be isolated from cells or tissue sources by an appropriate purification scheme using standard protein purification techniques. In another embodiment, ORFX proteins are produced by recombinant DNA techniques. Alternative to recombinant expression, a ORFX

protein or polypeptide can be synthesized chemically using standard peptide synthesis techniques.

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An "isolated" or "purified" protein or biologically active portion thereof is substantially free of cellular material or other contaminating proteins from the cell or tissue source from which the ORFX protein is derived, or substantially free from chemical precursors or other chemicals when chemically synthesized. The language "substantially free of cellular material" includes preparations of ORFX protein in which the protein is separated from cellular components of the cells from which it is isolated or recombinantly produced. In one embodiment, the language "substantially free of cellular material" includes preparations of ORFX protein having less than about 30% (by dry weight) of non-ORFX protein (also referred to herein as a "contaminating protein"), more preferably less than about 20% of non-ORFX protein, still more preferably less than about 10% of non-ORFX protein, and most preferably less than about 5% non-ORFX protein. When the ORFX protein or biologically active portion thereof is recombinantly produced, it is also preferably substantially free of culture medium, *i.e.*, culture medium represents less than about 20%, more preferably less than about 10%, and most preferably less than about 5% of the volume of the protein preparation.

The language "substantially free of chemical precursors or other chemicals" includes preparations of ORFX protein in which the protein is separated from chemical precursors or other chemicals that are involved in the synthesis of the protein. In one embodiment, the language "substantially free of chemical precursors or other chemicals" includes preparations of ORFX protein having less than about 30% (by dry weight) of chemical precursors or non-ORFX chemicals, more preferably less than about 20% chemical precursors or non-ORFX chemicals, still more preferably less than about 10% chemical precursors or non-ORFX chemicals, and most preferably less than about 5% chemical precursors or non-ORFX chemicals.

Biologically active portions of a ORFX protein include peptides comprising amino acid sequences sufficiently homologous to or derived from the amino acid sequence of the ORFX protein, e.g., the amino acid sequence shown in SEQ ID NO:2 that include fewer amino acids than the full length ORFX proteins, and exhibit at least one activity of a ORFX protein. Typically, biologically active portions comprise a domain or motif with at least one activity of the ORFX protein. A biologically active portion of a ORFX protein can be a polypeptide which is, for example, 10, 25, 50, 100 or more amino acids in length.

A biologically active portion of a ORFX protein of the present invention may contain at least one of the above-identified domains conserved between the FGF family of proteins. Moreover, other biologically active portions, in which other regions of the protein are deleted, can be prepared by recombinant techniques and evaluated for one or more of the functional activities of a native ORFX protein.

In an embodiment, the ORFX protein has an amino acid sequence shown in any of SEQ ID NO:2n (wherein n = 1 to 3161). In other embodiments, the ORFX protein is substantially homologous to any of SEQ ID NO:2n (wherein n = 1 to 3161) and retains the functional activity of the protein of any of SEQ ID NO:2n (wherein n = 1 to 3161), yet differs in amino acid sequence due to natural allelic variation or mutagenesis, as described in detail below. Accordingly, in another embodiment, the ORFX protein is a protein that comprises an amino acid sequence at least about 45% homologous, and more preferably about 55, 65, 70, 75, 80, 85, 90, 95, 98 or even 99% homologous to the amino acid sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161) and retains the functional activity of the ORFX proteins of the corresponding polypeptide having the sequence of SEQ ID NO:2n (wherein n = 1 to 3161).

Determining homology between two or more sequences

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To determine the percent homology of two amino acid sequences or of two nucleic acids, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in either of the sequences being compared for optimal alignment between the sequences). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are homologous at that position (i.e., as used herein amino acid or nucleic acid "homology" is equivalent to amino acid or nucleic acid "identity").

The nucleic acid sequence homology may be determined as the degree of identity between two sequences. The homology may be determined using computer programs known in the art, such as GAP software provided in the GCG program package. See, *Needleman and Wunsch* 1970 *J Mol Biol* 48: 443-453. Using GCG GAP software with the following settings for nucleic acid sequence comparison: GAP creation penalty of 5.0 and GAP extension penalty of 0.3, the coding region of the analogous nucleic acid sequences referred to above exhibits a

degree of identity preferably of at least 70%, 75%, 80%, 85%, 90%, 95%, 98%, or 99%, with the CDS (encoding) part of the DNA sequence shown in SEQ ID NO:2n-1 (wherein n = 1 to 3161).

The term "sequence identity" refers to the degree to which two polynucleotide or polypeptide sequences are identical on a residue-by-residue basis over a particular region of comparison. The term "percentage of sequence identity" is calculated by comparing two optimally aligned sequences over that region of comparison, determining the number of positions at which the identical nucleic acid base (e.g., A, T, C, G, U, or I, in the case of nucleic acids) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the region of comparison (i.e., the window size), and multiplying the result by 100 to yield the percentage of sequence identity. The term "substantial identity" as used herein denotes a characteristic of a polynucleotide sequence, wherein the polynucleotide comprises a sequence that has at least 80 percent sequence identity, preferably at least 85 percent identity and often 90 to 95 percent sequence identity, more usually at least 99 percent sequence identity as compared to a reference sequence over a comparison region. The term "percentage of positive residues" is calculated by comparing two optimally aligned sequences over that region of comparison, determining the number of positions at which the identical and conservative amino acid substitutions, as defined above, occur in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the region of comparison (i.e., the window size), and multiplying the result by 100 to yield the percentage of positive residues.

Chimeric and fusion proteins

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The invention also provides ORFX chimeric or fusion proteins. As used herein, a ORFX "chimeric protein" or "fusion protein" includes a ORFX polypeptide operatively linked to a non-ORFX polypeptide. A "ORFX polypeptide" refers to a polypeptide having an amino acid sequence corresponding to ORFX, whereas a "non-ORFX polypeptide" refers to a polypeptide having an amino acid sequence corresponding to a protein that is not substantially homologous to the ORFX protein, e.g., a protein that is different from the ORFX protein and that is derived from the same or a different organism. Within a ORFX fusion protein the ORFX polypeptide can correspond to all or a portion of a ORFX protein. In one embodiment, a ORFX fusion protein comprises at least one biologically active portion of a ORFX protein. In another embodiment, a ORFX fusion protein comprises at least two biologically active portions of a

ORFX protein. Within the fusion protein, the term "operatively linked" is intended to indicate that the ORFX polypeptide and the non-ORFX polypeptide are fused in-frame to each other. The non-ORFX polypeptide can be fused to the N-terminus or C-terminus of the ORFX polypeptide.

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For example, in one embodiment a ORFX fusion protein comprises a ORFX polypeptide operably linked to the extracellular domain of a second protein. Such fusion proteins can be further utilized in screening assays for compounds that modulate ORFX activity (such assays are described in detail below).

In another embodiment, the fusion protein is a GST-ORFX fusion protein in which the ORFX sequences are fused to the C-terminus of the GST (i.e., glutathione S-transferase) sequences. Such fusion proteins can facilitate the purification of recombinant ORFX.

In yet another embodiment, the fusion protein is a ORFX protein containing a heterologous signal sequence at its N-terminus. For example, the native ORFX signal sequence can be removed and replaced with a signal sequence from another protein. In certain host cells (e.g., mammalian host cells), expression and/or secretion of ORFX can be increased through use of a heterologous signal sequence.

In another embodiment, the fusion protein is a ORFX-immunoglobulin fusion protein in which the ORFX sequences comprising one or more domains are fused to sequences derived from a member of the immunoglobulin protein family. The ORFX-immunoglobulin fusion proteins of the invention can be incorporated into pharmaceutical compositions and administered to a subject to inhibit an interaction between a ORFX ligand and a ORFX protein on the surface of a cell, to thereby suppress ORFX-mediated signal transduction *in vivo*. In one nonlimiting example, a contemplated ORFX ligand of the invention is an ORFX receptor. The ORFX-immunoglobulin fusion proteins can be used to modulate the bioavailability of a ORFX cognate ligand. Inhibition of the ORFX ligand/ORFX interaction may be useful therapeutically for both the treatment of proliferative and differentiative disorders, as well as modulating (e.g., promoting or inhibiting) cell survival. Moreover, the ORFX-immunoglobulin fusion proteins of the invention can be used as immunogens to produce anti-ORFX antibodies in a subject, to purify ORFX ligands, and in screening assays to identify molecules that inhibit the interaction of ORFX with a ORFX ligand.

A ORFX chimeric or fusion protein of the invention can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with conventional techniques, e.g., by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers that give rise to complementary overhangs between two consecutive gene fragments that can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for example, Ausubel et al. (eds.) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). A ORFX-encoding nucleic acid can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the ORFX protein.

ORFX agonists and antagonists

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The present invention also pertains to variants of the ORFX proteins that function as either ORFX agonists (mimetics) or as ORFX antagonists. Variants of the ORFX protein can be generated by mutagenesis, e.g., discrete point mutation or truncation of the ORFX protein. An agonist of the ORFX protein can retain substantially the same, or a subset of, the biological activities of the naturally occurring form of the ORFX protein. An antagonist of the ORFX protein can inhibit one or more of the activities of the naturally occurring form of the ORFX protein by, for example, competitively binding to a downstream or upstream member of a cellular signaling cascade which includes the ORFX protein. Thus, specific biological effects can be elicited by treatment with a variant of limited function. In one embodiment, treatment of a subject with a variant having a subset of the biological activities of the naturally occurring form of the protein has fewer side effects in a subject relative to treatment with the naturally occurring form of the ORFX proteins.

Variants of the ORFX protein that function as either ORFX agonists (mimetics) or as ORFX antagonists can be identified by screening combinatorial libraries of mutants, e.g., truncation mutants, of the ORFX protein for ORFX protein agonist or antagonist activity. In one

embodiment, a variegated library of ORFX variants is generated by combinatorial mutagenesis at the nucleic acid level and is encoded by a variegated gene library. A variegated library of ORFX variants can be produced by, for example, enzymatically ligating a mixture of synthetic oligonucleotides into gene sequences such that a degenerate set of potential ORFX sequences is expressible as individual polypeptides, or alternatively, as a set of larger fusion proteins (e.g., for phage display) containing the set of ORFX sequences therein. There are a variety of methods which can be used to produce libraries of potential ORFX variants from a degenerate oligonucleotide sequence. Chemical synthesis of a degenerate gene sequence can be performed in an automatic DNA synthesizer, and the synthetic gene then ligated into an appropriate expression vector. Use of a degenerate set of genes allows for the provision, in one mixture, of all of the sequences encoding the desired set of potential ORFX sequences. Methods for synthesizing degenerate oligonucleotides are known in the art (see, e.g., Narang (1983) Tetrahedron 39:3; Itakura et al. (1984) Annu Rev Biochem 53:323; Itakura et al. (1984) Science 198:1056; Ike et al. (1983) Nucl Acid Res 11:477.

Polypeptide libraries

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In addition, libraries of fragments of the ORFX protein coding sequence can be used to generate a variegated population of ORFX fragments for screening and subsequent selection of variants of a ORFX protein. In one embodiment, a library of coding sequence fragments can be generated by treating a double stranded PCR fragment of a ORFX coding sequence with a nuclease under conditions wherein nicking occurs only about once per molecule, denaturing the double stranded DNA, renaturing the DNA to form double stranded DNA that can include sense/antisense pairs from different nicked products, removing single stranded portions from reformed duplexes by treatment with S1 nuclease, and ligating the resulting fragment library into an expression vector. By this method, an expression library can be derived which encodes N-terminal and internal fragments of various sizes of the ORFX protein.

Several techniques are known in the art for screening gene products of combinatorial libraries made by point mutations or truncation, and for screening cDNA libraries for gene products having a selected property. Such techniques are adaptable for rapid screening of the gene libraries generated by the combinatorial mutagenesis of ORFX proteins. The most widely used techniques, which are amenable to high throughput analysis, for screening large gene libraries typically include cloning the gene library into replicable expression vectors,

transforming appropriate cells with the resulting library of vectors, and expressing the combinatorial genes under conditions in which detection of a desired activity facilitates isolation of the vector encoding the gene whose product was detected. Recrusive ensemble mutagenesis (REM), a new technique that enhances the frequency of functional mutants in the libraries, can be used in combination with the screening assays to identify ORFX variants (Arkin and Yourvan (1992) PNAS 89:7811-7815; Delgrave *et al.* (1993) Protein Engineering 6:327-331).

Anti-ORFX Antibodies

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The invention further encompasses antibodies and antibody fragments, such as F_{ab} or $(F_{ab})_2$, that bind immunospecifically to any of the proteins of the invention.

An isolated ORFX protein, or a portion or fragment thereof, can be used as an immunogen to generate antibodies that bind ORFX using standard techniques for polyclonal and monoclonal antibody preparation. Full-length ORFX protein can be used. Alternatively, the invention provides antigenic peptide fragments of ORFX for use as immunogens. The antigenic peptide of ORFX comprises at least 4 amino acid residues of the amino acid sequence shown in any of SEQ ID NO:2n (wherein n = 1 to 3161). The antigenic peptide encompasses an epitope of ORFX such that an antibody raised against the peptide forms a specific immune complex with ORFX. The antigenic peptide may comprise at least 6 aa residues, at least 8 aa residues, at least 10 aa residues, at least 15 aa residues, at least 20 aa residues, or at least 30 aa residues. In one embodiment of the invention, the antigenic peptide comprises a polypeptide comprising at least 6 contiguous amino acids of any of SEQ ID NO:2n (wherein n = 1 to 3161).

In an embodiment of the invention, epitopes encompassed by the antigenic peptide are regions of ORFX that are located on the surface of the protein, e.g., hydrophilic regions. As a means for targeting antibody production, hydropathy plots showing regions of hydrophilicity and hydrophobicity may be generated by any method well known in the art, including, for example, the Kyte Doolittle or the Hopp Woods methods, either with or without Fourier transformation. See, e.g., Hopp and Woods, 1981, Proc. Nat. Acad. Sci. USA 78: 3824-3828; Kyte and Doolittle 1982, J. Mol. Biol. 157: 105-142, each incorporated herein by reference in their entirety.

As disclosed herein, an ORFX protein sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161), or derivatives, fragments, analogs or homologs thereof, may be utilized as immunogens in the generation of antibodies that immunospecifically-bind these protein components. The term "antibody" as used herein refers to immunoglobulin molecules and

immunologically active portions of immunoglobulin molecules, i.e., molecules that contain an antigen binding site that specifically binds (immunoreacts with) an antigen, such as ORFX. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain, F_{ab} and $F_{(ab)2}$ fragments, and an F_{ab} expression library. In a specific embodiment, antibodies to human ORFX proteins are disclosed. Various procedures known within the art may be used for the production of polyclonal or monoclonal antibodies to a ORFX protein sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161) or derivative, fragment, analog or homolog thereof. Some of these proteins are discussed below.

For the production of polyclonal antibodies, various suitable host animals (e.g., rabbit, goat, mouse or other mammal) may be immunized by injection with the native protein, or a synthetic variant thereof, or a derivative of the foregoing. An appropriate immunogenic preparation can contain, for example, recombinantly expressed ORFX protein or a chemically synthesized ORFX polypeptide. The preparation can further include an adjuvant. Various adjuvants used to increase the immunological response include, but are not limited to, Freund's (complete and incomplete), mineral gels (e.g., aluminum hydroxide), surface active substances (e.g., lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, etc.), human adjuvants such as Bacille Calmette-Guerin and Corynebacterium parvum, or similar immunostimulatory agents. If desired, the antibody molecules directed against ORFX can be isolated from the mammal (e.g., from the blood) and further purified by well known techniques, such as protein A chromatography to obtain the IgG fraction.

The term "monoclonal antibody" or "monoclonal antibody composition", as used herein, refers to a population of antibody molecules that contain only one species of an antigen binding site capable of immunoreacting with a particular epitope of ORFX. A monoclonal antibody composition thus typically displays a single binding affinity for a particular ORFX protein with which it immunoreacts. For preparation of monoclonal antibodies directed towards a particular ORFX protein, or derivatives, fragments, analogs or homologs thereof, any technique that provides for the production of antibody molecules by continuous cell line culture may be utilized. Such techniques include, but are not limited to, the hybridoma technique (see Kohler & Milstein, 1975 *Nature* 256: 495-497); the trioma technique; the human B-cell hybridoma technique (see Kozbor, *et al.*, 1983 *Immunol Today* 4: 72) and the EBV hybridoma technique to produce human monoclonal antibodies (see Cole, *et al.*, 1985 In: MONOCLONAL ANTIBODIES AND

CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Human monoclonal antibodies may be utilized in the practice of the present invention and may be produced by using human hybridomas (see Cote, et al., 1983. Proc Natl Acad Sci USA 80: 2026-2030) or by transforming human B-cells with Epstein Barr Virus in vitro (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Each of the above citations are incorporated herein by reference in their entirety

According to the invention, techniques can be adapted for the production of single-chain antibodies specific to a ORFX protein (see e.g., U.S. Patent No. 4,946,778). In addition, methods can be adapted for the construction of F_{ab} expression libraries (see e.g., Huse, et al., 1989 Science 246: 1275-1281) to allow rapid and effective identification of monoclonal F_{ab} fragments with the desired specificity for a ORFX protein or derivatives, fragments, analogs or homologs thereof. Non-human antibodies can be "humanized" by techniques well known in the art. See e.g., U.S. Patent No. 5,225,539. Each of the above citations are incorporated herein by reference. Antibody fragments that contain the idiotypes to a ORFX protein may be produced by techniques known in the art including, but not limited to: (i) an $F_{(ab)2}$ fragment produced by pepsin digestion of an antibody molecule; (ii) an F_{ab} fragment generated by reducing the disulfide bridges of an $F_{(ab)2}$ fragment; (iii) an F_{ab} fragment generated by the treatment of the antibody molecule with papain and a reducing agent and (iv) F_{v} fragments.

Additionally, recombinant anti-ORFX antibodies, such as chimeric and humanized monoclonal antibodies, comprising both human and non-human portions, which can be made using standard recombinant DNA techniques, are within the scope of the invention. Such chimeric and humanized monoclonal antibodies can be produced by recombinant DNA techniques known in the art, for example using methods described in PCT International Application No. PCT/US86/02269; European Patent Application No. 184,187; European Patent Application No. 171,496; European Patent Application No. 173,494; PCT International Publication No. WO 86/01533; U.S. Pat. No. 4,816,567; European Patent Application No. 125,023; Better et al.(1988) Science 240:1041-1043; Liu et al. (1987) PNAS 84:3439-3443; Liu et al. (1987) J Immunol. 139:3521-3526; Sun et al. (1987) PNAS 84:214-218; Nishimura et al. (1987) Cancer Res 47:999-1005; Wood et al. (1985) Nature 314:446-449; Shaw et al. (1988), J. Natl Cancer Inst 80:1553-1559); Morrison(1985) Science 229:1202-1207; Oi et al. (1986) BioTechniques 4:214; U.S. Pat. No. 5,225,539; Jones et al. (1986) Nature 321:552-525;

Verhoeyan et al. (1988) Science 239:1534; and Beidler et al. (1988) J Immunol 141:4053-4060. Each of the above citations are incorporated herein by reference.

In one embodiment, methods for the screening of antibodies that possess the desired specificity include, but are not limited to, enzyme-linked immunosorbent assay (ELISA) and other immunologically-mediated techniques known within the art. In a specific embodiment, selection of antibodies that are specific to a particular domain of a ORFX protein is facilitated by generation of hybridomas that bind to the fragment of a ORFX protein possessing such a domain. Antibodies that are specific for one or more domains within a ORFX protein, e.g., the domain spanning the first fifty amino-terminal residues specific to ORFX when compared to FGF-9, or derivatives, fragments, analogs or homologs thereof, are also provided herein.

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Anti-ORFX antibodies may be used in methods known within the art relating to the localization and/or quantitation of a ORFX protein (e.g., for use in measuring levels of the ORFX protein within appropriate physiological samples, for use in diagnostic methods, for use in imaging the protein, and the like). In a given embodiment, antibodies for ORFX proteins, or derivatives, fragments, analogs or homologs thereof, that contain the antibody derived binding domain, are utilized as pharmacologically-active compounds [hereinafter "Therapeutics"].

An anti-ORFX antibody (e.g., monoclonal antibody) can be used to isolate ORFX by standard techniques, such as affinity chromatography or immunoprecipitation. An anti-ORFX antibody can facilitate the purification of natural ORFX from cells and of recombinantly produced ORFX expressed in host cells. Moreover, an anti-ORFX antibody can be used to detect ORFX protein (e.g., in a cellular lysate or cell supernatant) in order to evaluate the abundance and pattern of expression of the ORFX protein. Anti-ORFX antibodies can be used diagnostically to monitor protein levels in tissue as part of a clinical testing procedure, e.g., to, for example, determine the efficacy of a given treatment regimen. Detection can be facilitated by coupling (i.e., physically linking) the antibody to a detectable substance. Examples of detectable substances include various enzymes, prosthetic groups, fluorescent materials, luminescent materials, bioluminescent materials, and radioactive materials. Examples of suitable enzymes include horseradish peroxidase, alkaline phosphatase, β-galactosidase, or acetylcholinesterase; examples of suitable prosthetic group complexes include streptavidin/biotin and avidin/biotin; examples of suitable fluorescent materials include umbelliferone, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dansyl chloride or

phycoerythrin; an example of a luminescent material includes luminol; examples of bioluminescent materials include luciferase, luciferin, and aequorin, and examples of suitable radioactive material include ¹²⁵I, ¹³¹I, ³⁵S or ³H.

ORFX Recombinant Vectors and Host Cells

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Another aspect of the invention pertains to vectors, preferably expression vectors, containing a nucleic acid encoding ORFX protein, or derivatives, fragments, analogs or homologs thereof. As used herein, the term "vector" refers to a nucleic acid molecule capable of transporting another nucleic acid to which it has been linked. One type of vector is a "plasmid", which refers to a circular double stranded DNA loop into which additional DNA segments can be ligated. Another type of vector is a viral vector, wherein additional DNA segments can be ligated into the viral genome. Certain vectors are capable of autonomous replication in a host cell into which they are introduced (e.g., bacterial vectors having a bacterial origin of replication and episomal mammalian vectors). Other vectors (e.g., non-episomal mammalian vectors) are integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome. Moreover, certain vectors are capable of directing the expression of genes to which they are operatively linked. Such vectors are referred to herein as "expression vectors". In general, expression vectors of utility in recombinant DNA techniques are often in the form of plasmids. In the present specification, "plasmid" and "vector" can be used interchangeably as the plasmid is the most commonly used form of vector. However, the invention is intended to include such other forms of expression vectors, such as viral vectors (e.g., replication defective retroviruses, adenoviruses and adeno-associated viruses), which serve equivalent functions.

The recombinant expression vectors of the invention comprise a nucleic acid of the invention in a form suitable for expression of the nucleic acid in a host cell, which means that the recombinant expression vectors include one or more regulatory sequences, selected on the basis of the host cells to be used for expression, that is operatively linked to the nucleic acid sequence to be expressed. Within a recombinant expression vector, "operably linked" is intended to mean that the nucleotide sequence of interest is linked to the regulatory sequence(s) in a manner that allows for expression of the nucleotide sequence (e.g., in an in vitro transcription/translation system or in a host cell when the vector is introduced into the host cell). The term "regulatory sequence" is intended to includes promoters, enhancers and other expression control elements

(e.g., polyadenylation signals). Such regulatory sequences are described, for example, in Goeddel; GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990). Regulatory sequences include those that direct constitutive expression of a nucleotide sequence in many types of host cell and those that direct expression of the nucleotide sequence only in certain host cells (e.g., tissue-specific regulatory sequences). It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host cell to be transformed, the level of expression of protein desired, etc. The expression vectors of the invention can be introduced into host cells to thereby produce proteins or peptides, including fusion proteins or peptides, encoded by nucleic acids as described herein (e.g., ORFX proteins, mutant forms of ORFX, fusion proteins, etc.).

The recombinant expression vectors of the invention can be designed for expression of ORFX in prokaryotic or eukaryotic cells. For example, ORFX can be expressed in bacterial cells such as *E. coli*, insect cells (using baculovirus expression vectors) yeast cells or mammalian cells. Suitable host cells are discussed further in Goeddel, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990). Alternatively, the recombinant expression vector can be transcribed and translated *in vitro*, for example using T7 promoter regulatory sequences and T7 polymerase.

Expression of proteins in prokaryotes is most often carried out in *E. coli* with vectors containing constitutive or inducible promoters directing the expression of either fusion or non-fusion proteins. Fusion vectors add a number of amino acids to a protein encoded therein, usually to the amino terminus of the recombinant protein. Such fusion vectors typically serve three purposes: (1) to increase expression of recombinant protein; (2) to increase the solubility of the recombinant protein; and (3) to aid in the purification of the recombinant protein by acting as a ligand in affinity purification. Often, in fusion expression vectors, a proteolytic cleavage site is introduced at the junction of the fusion moiety and the recombinant protein to enable separation of the recombinant protein from the fusion moiety subsequent to purification of the fusion protein. Such enzymes, and their cognate recognition sequences, include Factor Xa, thrombin and enterokinase. Typical fusion expression vectors include pGEX (Pharmacia Biotech Inc; Smith and Johnson (1988) *Gene* 67:31-40), pMAL (New England Biolabs, Beverly, Mass.) and pRIT5 (Pharmacia, Piscataway, N.J.) that fuse glutathione S-transferase (GST), maltose E binding protein, or protein A, respectively, to the target recombinant protein.

Examples of suitable inducible non-fusion *E. coli* expression vectors include pTrc (Amrann *et al.*, (1988) *Gene* 69:301-315) and pET 11d (Studier *et al.*, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990) 60-89).

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One strategy to maximize recombinant protein expression in *E. coli* is to express the protein in a host bacteria with an impaired capacity to proteolytically cleave the recombinant protein. See, Gottesman, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990) 119-128. Another strategy is to alter the nucleic acid sequence of the nucleic acid to be inserted into an expression vector so that the individual codons for each amino acid are those preferentially utilized in *E. coli* (Wada *et al.*, (1992) *Nucleic Acids Res.* 20:2111-2118). Such alteration of nucleic acid sequences of the invention can be carried out by standard DNA synthesis techniques.

In another embodiment, the ORFX expression vector is a yeast expression vector. Examples of vectors for expression in yeast *S. cerivisae* include pYepSec1 (Baldari, *et al.*, (1987) *EMBO J* 6:229-234), pMFa (Kurjan and Herskowitz, (1982) *Cell* 30:933-943), pJRY88 (Schultz *et al.*, (1987) *Gene* 54:113-123), pYES2 (Invitrogen Corporation, San Diego, Calif.), and picZ (InVitrogen Corp, San Diego, Calif.).

Alternatively, ORFX can be expressed in insect cells using baculovirus expression vectors. Baculovirus vectors available for expression of proteins in cultured insect cells (e.g., SF9 cells) include the pAc series (Smith et al. (1983) Mol Cell Biol 3:2156-2165) and the pVL series (Lucklow and Summers (1989) Virology 170:31-39).

In yet another embodiment, a nucleic acid of the invention is expressed in mammalian cells using a mammalian expression vector. Examples of mammalian expression vectors include pCDM8 (Seed (1987) Nature 329:840) and pMT2PC (Kaufman et al. (1987) EMBO J 6: 187-195). When used in mammalian cells, the expression vector's control functions are often provided by viral regulatory elements. For example, commonly used promoters are derived from polyoma, Adenovirus 2, cytomegalovirus and Simian Virus 40. For other suitable expression systems for both prokaryotic and eukaryotic cells. See, e.g., Chapters 16 and 17 of Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL. 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989.

In another embodiment, the recombinant mammalian expression vector is capable of directing expression of the nucleic acid preferentially in a particular cell type (e.g., tissue-specific regulatory elements are used to express the nucleic acid). Tissue-specific regulatory elements are known in the art. Non-limiting examples of suitable tissue-specific promoters include the albumin promoter (liver-specific; Pinkert et al. (1987) Genes Dev 1:268-277), lymphoid-specific promoters (Calame and Eaton (1988) Adv Immunol 43:235-275), in particular promoters of T cell receptors (Winoto and Baltimore (1989) EMBO J 8:729-733) and immunoglobulins (Banerji et al. (1983) Cell 33:729-740; Queen and Baltimore (1983) Cell 33:741-748), neuron-specific promoters (e.g., the neurofilament promoter; Byrne and Ruddle (1989) PNAS 86:5473-5477), pancreas-specific promoters (Edlund et al. (1985) Science 230:912-916), and mammary gland-specific promoters (e.g., milk whey promoter; U.S. Pat. No. 4,873,316 and European Application Publication No. 264,166). Developmentally-regulated promoters are also encompassed, e.g., the murine hox promoters (Kessel and Gruss (1990) Science 249:374-379) and the α-fetoprotein promoter (Campes and Tilghman (1989) Genes Dev 3:537-546).

The invention further provides a recombinant expression vector comprising a DNA molecule of the invention cloned into the expression vector in an antisense orientation. That is, the DNA molecule is operatively linked to a regulatory sequence in a manner that allows for expression (by transcription of the DNA molecule) of an RNA molecule that is antisense to ORFX mRNA. Regulatory sequences operatively linked to a nucleic acid cloned in the antisense orientation can be chosen that direct the continuous expression of the antisense RNA molecule in a variety of cell types, for instance viral promoters and/or enhancers, or regulatory sequences can be chosen that direct constitutive, tissue specific or cell type specific expression of antisense RNA. The antisense expression vector can be in the form of a recombinant plasmid, phagemid or attenuated virus in which antisense nucleic acids are produced under the control of a high efficiency regulatory region, the activity of which can be determined by the cell type into which the vector is introduced. For a discussion of the regulation of gene expression using antisense genes see Weintraub *et al.*, "Antisense RNA as a molecular tool for genetic analysis," Reviews--Trends in Genetics, Vol. 1(1) 1986.

Another aspect of the invention pertains to host cells into which a recombinant expression vector of the invention has been introduced. The terms "host cell" and "recombinant

host cell" are used interchangeably herein. It is understood that such terms refer not only to the particular subject cell but to the progeny or potential progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not, in fact, be identical to the parent cell, but are still included within the scope of the term as used herein.

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A host cell can be any prokaryotic or eukaryotic cell. For example, ORFX protein can be expressed in bacterial cells such as *E. coli*, insect cells, yeast or mammalian cells (such as Chinese hamster ovary cells (CHO) or COS cells). Other suitable host cells are known to those skilled in the art.

Vector DNA can be introduced into prokaryotic or eukaryotic cells via conventional transformation or transfection techniques. As used herein, the terms "transformation" and "transfection" are intended to refer to a variety of art-recognized techniques for introducing foreign nucleic acid (e.g., DNA) into a host cell, including calcium phosphate or calcium chloride co-precipitation, DEAE-dextran-mediated transfection, lipofection, or electroporation. Suitable methods for transforming or transfecting host cells can be found in Sambrook, et al. (MOLECULAR CLONING: A LABORATORY MANUAL. 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989), and other laboratory manuals.

For stable transfection of mammalian cells, it is known that, depending upon the expression vector and transfection technique used, only a small fraction of cells may integrate the foreign DNA into their genome. In order to identify and select these integrants, a gene that encodes a selectable marker (e.g., resistance to antibiotics) is generally introduced into the host cells along with the gene of interest. Various selectable markers include those that confer resistance to drugs, such as G418, hygromycin and methotrexate. Nucleic acid encoding a selectable marker can be introduced into a host cell on the same vector as that encoding ORFX or can be introduced on a separate vector. Cells stably transfected with the introduced nucleic acid can be identified by drug selection (e.g., cells that have incorporated the selectable marker gene will survive, while the other cells die).

A host cell of the invention, such as a prokaryotic or eukaryotic host cell in culture, can be used to produce (i.e., express) ORFX protein. Accordingly, the invention further provides methods for producing ORFX protein using the host cells of the invention. In one embodiment,

the method comprises culturing the host cell of invention (into which a recombinant expression vector encoding ORFX has been introduced) in a suitable medium such that ORFX protein is produced. In another embodiment, the method further comprises isolating ORFX from the medium or the host cell.

Transgenic animals

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The host cells of the invention can also be used to produce nonhuman transgenic animals. For example, in one embodiment, a host cell of the invention is a fertilized oocyte or an embryonic stem cell into which ORFX-coding sequences have been introduced. Such host cells can then be used to create non-human transgenic animals in which exogenous ORFX sequences have been introduced into their genome or homologous recombinant animals in which endogenous ORFX sequences have been altered. Such animals are useful for studying the function and/or activity of ORFX and for identifying and/or evaluating modulators of ORFX activity. As used herein, a "transgenic animal" is a non-human animal, preferably a mammal, more preferably a rodent such as a rat or mouse, in which one or more of the cells of the animal includes a transgene. Other examples of transgenic animals include non-human primates, sheep, dogs, cows, goats, chickens, amphibians, etc. A transgene is exogenous DNA that is integrated into the genome of a cell from which a transgenic animal develops and that remains in the genome of the mature animal, thereby directing the expression of an encoded gene product in one or more cell types or tissues of the transgenic animal. As used herein, a "homologous recombinant animal" is a non-human animal, preferably a mammal, more preferably a mouse, in which an endogenous ORFX gene has been altered by homologous recombination between the endogenous gene and an exogenous DNA molecule introduced into a cell of the animal, e.g., an embryonic cell of the animal, prior to development of the animal.

A transgenic animal of the invention can be created by introducing ORFX-encoding nucleic acid into the male pronuclei of a fertilized oocyte, e.g., by microinjection, retroviral infection, and allowing the oocyte to develop in a pseudopregnant female foster animal. The human ORFX DNA sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161) can be introduced as a transgene into the genome of a non-human animal. Alternatively, a nonhuman homologue of the human ORFX gene, such as a mouse ORFX gene, can be isolated based on hybridization to the human ORFX cDNA (described further above) and used as a transgene. Intronic sequences and polyadenylation signals can also be included in the transgene to increase the efficiency of

expression of the transgene. A tissue-specific regulatory sequence(s) can be operably linked to the ORFX transgene to direct expression of ORFX protein to particular cells. Methods for generating transgenic animals via embryo manipulation and microinjection, particularly animals such as mice, have become conventional in the art and are described, for example, in U.S. Pat. Nos. 4,736,866; 4,870,009; and 4,873,191; and Hogan 1986, In: Manipulating the Mouse Embryo, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. Similar methods are used for production of other transgenic animals. A transgenic founder animal can be identified based upon the presence of the ORFX transgene in its genome and/or expression of ORFX mRNA in tissues or cells of the animals. A transgenic founder animal can then be used to breed additional animals carrying the transgene. Moreover, transgenic animals carrying a transgene encoding ORFX can further be bred to other transgenic animals carrying other transgenes.

To create a homologous recombinant animal, a vector is prepared which contains at least a portion of a ORFX gene into which a deletion, addition or substitution has been introduced to thereby alter, e.g., functionally disrupt, the ORFX gene. The ORFX gene can be a human gene (e.g., SEQ ID NO:2n-1 (wherein n=1 to 3161)), but more preferably, is a non-human homologue of a human ORFX gene. For example, a mouse homologue of human ORFX gene of SEQ ID NO:2n-1 (wherein n=1 to 3161) can be used to construct a homologous recombination vector suitable for altering an endogenous ORFX gene in the mouse genome. In one embodiment, the vector is designed such that, upon homologous recombination, the endogenous ORFX gene is functionally disrupted (i.e., no longer encodes a functional protein; also referred to as a "knock out" vector).

Alternatively, the vector can be designed such that, upon homologous recombination, the endogenous ORFX gene is mutated or otherwise altered but still encodes functional protein (e.g., the upstream regulatory region can be altered to thereby alter the expression of the endogenous ORFX protein). In the homologous recombination vector, the altered portion of the ORFX gene is flanked at its 5' and 3' ends by additional nucleic acid of the ORFX gene to allow for homologous recombination to occur between the exogenous ORFX gene carried by the vector and an endogenous ORFX gene in an embryonic stem cell. The additional flanking ORFX nucleic acid is of sufficient length for successful homologous recombination with the endogenous gene. Typically, several kilobases of flanking DNA (both at the 5' and 3' ends) are included in the vector. See e.g., Thomas et al. (1987) Cell 51:503 for a description of

homologous recombination vectors. The vector is introduced into an embryonic stem cell line (e.g., by electroporation) and cells in which the introduced ORFX gene has homologously recombined with the endogenous ORFX gene are selected (see e.g., Li et al. (1992) Cell 69:915).

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The selected cells are then injected into a blastocyst of an animal (e.g., a mouse) to form aggregation chimeras. See e.g., Bradley 1987, In: TERATOCARCINOMAS AND EMBRYONIC STEM CELLS: A PRACTICAL APPROACH, Robertson, ed. IRL, Oxford, pp. 113-152. A chimeric embryo can then be implanted into a suitable pseudopregnant female foster animal and the embryo brought to term. Progeny harboring the homologously recombined DNA in their germ cells can be used to breed animals in which all cells of the animal contain the homologously recombined DNA by germline transmission of the transgene. Methods for constructing homologous recombination vectors and homologous recombinant animals are described further in Bradley (1991) Curr Opin Biotechnol 2:823-829; PCT International Publication Nos.: WO 90/11354; WO 91/01140; WO 92/0968; and WO 93/04169.

In another embodiment, transgenic non-humans animals can be produced that contain selected systems that allow for regulated expression of the transgene. One example of such a system is the cre/loxP recombinase system of bacteriophage P1. For a description of the cre/loxP recombinase system, see, e.g., Lakso et al. (1992) PNAS 89:6232-6236. Another example of a recombinase system is the FLP recombinase system of Saccharomyces cerevisiae (O'Gorman et al. (1991) Science 251:1351-1355. If a cre/loxP recombinase system is used to regulate expression of the transgene, animals containing transgenes encoding both the Cre recombinase and a selected protein are required. Such animals can be provided through the construction of "double" transgenic animals, e.g., by mating two transgenic animals, one containing a transgene encoding a selected protein and the other containing a transgene encoding a recombinase.

Clones of the non-human transgenic animals described herein can also be produced according to the methods described in Wilmut et al. (1997) Nature 385:810-813. In brief, a cell, e.g., a somatic cell, from the transgenic animal can be isolated and induced to exit the growth cycle and enter G₀ phase. The quiescent cell can then be fused, e.g., through the use of electrical pulses, to an enucleated oocyte from an animal of the same species from which the quiescent cell is isolated. The reconstructed oocyte is then cultured such that it develops to morula or blastocyte and then transferred to pseudopregnant female foster animal. The offspring borne of

this female foster animal will be a clone of the animal from which the cell, e.g., the somatic cell, is isolated.

Pharmaceutical Compositions

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The ORFX nucleic acid molecules, ORFX proteins, and anti-ORFX antibodies (also referred to herein as "active compounds") of the invention, and derivatives, fragments, analogs and homologs thereof, can be incorporated into pharmaceutical compositions suitable for administration. Such compositions typically comprise the nucleic acid molecule, protein, or antibody and a pharmaceutically acceptable carrier. As used herein, "pharmaceutically acceptable carrier" is intended to include any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like, compatible with pharmaceutical administration. Suitable carriers are described in the most recent edition of Remington's Pharmaceutical Sciences, a standard reference text in the field, which is incorporated herein by reference. Preferred examples of such carriers or diluents include, but are not limited to, water, saline, finger's solutions, dextrose solution, and 5% human serum albumin. Liposomes and non-aqueous vehicles such as fixed oils may also be used. The use of such media and agents for pharmaceutically active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active compound, use thereof in the compositions is contemplated. Supplementary active compounds can also be incorporated into the compositions.

A pharmaceutical composition of the invention is formulated to be compatible with its intended route of administration. Examples of routes of administration include parenteral, e.g., intravenous, intradermal, subcutaneous, oral (e.g., inhalation), transdermal (topical), transmucosal, and rectal administration. Solutions or suspensions used for parenteral, intradermal, or subcutaneous application can include the following components: a sterile diluent such as water for injection, saline solution, fixed oils, polyethylene glycols, glycerine, propylene glycol or other synthetic solvents; antibacterial agents such as benzyl alcohol or methyl parabens; antioxidants such as ascorbic acid or sodium bisulfite; chelating agents such as ethylenediaminetetraacetic acid; buffers such as acetates, citrates or phosphates, and agents for the adjustment of tonicity such as sodium chloride or dextrose. The pH can be adjusted with acids or bases, such as hydrochloric acid or sodium hydroxide. The parenteral preparation can be enclosed in ampoules, disposable syringes or multiple dose vials made of glass or plastic.

Pharmaceutical compositions suitable for injectable use include sterile aqueous solutions (where water soluble) or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersion. For intravenous administration, suitable carriers include physiological saline, bacteriostatic water, Cremophor EL™ (BASF, Parsippany, N.J.) or phosphate buffered saline (PBS). In all cases, the composition must be sterile and should be fluid to the extent that easy syringeability exists. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (for example, glycerol, propylene glycol, and liquid polyethylene glycol and the like), and suitable mixtures thereof. The proper fluidity can be maintained, for example, by the use of a coating such as lecithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants. Prevention of the action of microorganisms can be achieved by various antibacterial and antifungal agents, for example, parabens, chlorobutanol, phenol, ascorbic acid, thimerosal, and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars, polyalcohols such as manitol, sorbitol, sodium chloride in the composition. Prolonged absorption of the injectable compositions can be brought about by including in the composition an agent which delays absorption, for example, aluminum monostearate and gelatin.

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Sterile injectable solutions can be prepared by incorporating the active compound (e.g., a ORFX protein or anti-ORFX antibody) in the required amount in an appropriate solvent with one or a combination of ingredients enumerated above, as required, followed by filtered sterilization. Generally, dispersions are prepared by incorporating the active compound into a sterile vehicle that contains a basic dispersion medium and the required other ingredients from those enumerated above. In the case of sterile powders for the preparation of sterile injectable solutions, methods of preparation are vacuum drying and freeze-drying that yields a powder of the active ingredient plus any additional desired ingredient from a previously sterile-filtered solution thereof.

Oral compositions generally include an inert diluent or an edible carrier. They can be enclosed in gelatin capsules or compressed into tablets. For the purpose of oral therapeutic administration, the active compound can be incorporated with excipients and used in the form of tablets, troches, or capsules. Oral compositions can also be prepared using a fluid carrier for use

as a mouthwash, wherein the compound in the fluid carrier is applied orally and swished and expectorated or swallowed. Pharmaceutically compatible binding agents, and/or adjuvant materials can be included as part of the composition. The tablets, pills, capsules, troches and the like can contain any of the following ingredients, or compounds of a similar nature: a binder such as microcrystalline cellulose, gum tragacanth or gelatin; an excipient such as starch or lactose, a disintegrating agent such as alginic acid, Primogel, or corn starch; a lubricant such as magnesium stearate or Sterotes; a glidant such as colloidal silicon dioxide; a sweetening agent such as sucrose or saccharin; or a flavoring agent such as peppermint, methyl salicylate, or orange flavoring.

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For administration by inhalation, the compounds are delivered in the form of an aerosol spray from pressured container or dispenser which contains a suitable propellant, e.g., a gas such as carbon dioxide, or a nebulizer.

Systemic administration can also be by transmucosal or transdermal means. For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art, and include, for example, for transmucosal administration, detergents, bile salts, and fusidic acid derivatives. Transmucosal administration can be accomplished through the use of nasal sprays or suppositories. For transdermal administration, the active compounds are formulated into ointments, salves, gels, or creams as generally known in the art.

The compounds can also be prepared in the form of suppositories (e.g., with conventional suppository bases such as cocoa butter and other glycerides) or retention enemas for rectal delivery.

In one embodiment, the active compounds are prepared with carriers that will protect the compound against rapid elimination from the body, such as a controlled release formulation, including implants and microencapsulated delivery systems. Biodegradable, biocompatible polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, collagen, polyorthoesters, and polylactic acid. Methods for preparation of such formulations will be apparent to those skilled in the art. The materials can also be obtained commercially from Alza Corporation and Nova Pharmaceuticals, Inc. Liposomal suspensions (including liposomes targeted to infected cells with monoclonal antibodies to viral antigens) can also be used as pharmaceutically acceptable carriers. These can be prepared according to methods known to those skilled in the art, for example, as described in U.S. Pat. No. 4,522,811.

It is especially advantageous to formulate oral or parenteral compositions in dosage unit form for ease of administration and uniformity of dosage. Dosage unit form as used herein refers to physically discrete units suited as unitary dosages for the subject to be treated; each unit containing a predetermined quantity of active compound calculated to produce the desired therapeutic effect in association with the required pharmaceutical carrier. The specification for the dosage unit forms of the invention are dictated by and directly dependent on the unique characteristics of the active compound and the particular therapeutic effect to be achieved.

The nucleic acid molecules of the invention can be inserted into vectors and used as gene therapy vectors. Gene therapy vectors can be delivered to a subject by any of a number of routes, e.g., as described in U.S. Patent Nos. 5,703,055. Delivery can thus also include, e.g., intravenous injection, local administration (see U.S. Pat. No. 5,328,470) or stereotactic injection (see e.g., Chen et al. (1994) PNAS 91:3054-3057). The pharmaceutical preparation of the gene therapy vector can include the gene therapy vector in an acceptable diluent, or can comprise a slow release matrix in which the gene delivery vehicle is imbedded. Alternatively, where the complete gene delivery vector can be produced intact from recombinant cells, e.g., retroviral vectors, the pharmaceutical preparation can include one or more cells that produce the gene delivery system.

The pharmaceutical compositions can be included in a container, pack, or dispenser together with instructions for administration.

Additional Uses and Methods of the Invention

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The nucleic acid molecules, proteins, protein homologues, and antibodies described herein can be used in one or more of the following methods: (a) screening assays; (b) detection assays (e.g., chromosomal mapping, cell and tissue typing, forensic biology), (c) predictive medicine (e.g., diagnostic assays, prognostic assays, monitoring clinical trials, and pharmacogenomics); and (d) methods of treatment (e.g., therapeutic and prophylactic).

The isolated nucleic acid molecules of the invention can be used to express ORFX protein (e.g., via a recombinant expression vector in a host cell in gene therapy applications), to detect ORFX mRNA (e.g., in a biological sample) or a genetic lesion in a ORFX gene, and to modulate ORFX activity, as described further below. In addition, the ORFX proteins can be used to screen drugs or compounds that modulate the ORFX activity or expression as well as to treat disorders characterized by insufficient or excessive production of ORFX protein, for

example proliferative or differentiative disorders, or production of ORFX protein forms that have decreased or aberrant activity compared to ORFX wild type protein. In addition, the anti-ORFX antibodies of the invention can be used to detect and isolate ORFX proteins and modulate ORFX activity.

This invention further pertains to novel agents identified by the above described screening assays and uses thereof for treatments as described herein.

Screening Assays

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The invention provides a method (also referred to herein as a "screening assay") for identifying modulators, *i.e.*, candidate or test compounds or agents (*e.g.*, peptides, peptidomimetics, small molecules or other drugs) that bind to ORFX proteins or have a stimulatory or inhibitory effect on, for example, ORFX expression or ORFX activity.

In one embodiment, the invention provides assays for screening candidate or test compounds which bind to or modulate the activity of a ORFX protein or polypeptide or biologically active portion thereof. The test compounds of the present invention can be obtained using any of the numerous approaches in combinatorial library methods known in the art, including: biological libraries; spatially addressable parallel solid phase or solution phase libraries; synthetic library methods requiring deconvolution; the "one-bead one-compound" library method; and synthetic library methods using affinity chromatography selection. The biological library approach is limited to peptide libraries, while the other four approaches are applicable to peptide, non-peptide oligomer or small molecule libraries of compounds (Lam (1997) Anticancer Drug Des 12:145).

Examples of methods for the synthesis of molecular libraries can be found in the art, for example in: DeWitt et al. (1993) Proc Natl Acad Sci U.S.A. 90:6909; Erb et al. (1994) Proc Natl Acad Sci U.S.A. 91:11422; Zuckermann et al. (1994) J Med Chem 37:2678; Cho et al. (1993) Science 261:1303; Carrell et al. (1994) Angew Chem Int Ed Engl 33:2059; Carell et al. (1994) Angew Chem Int Ed Engl 33:2061; and Gallop et al. (1994) J Med Chem 37:1233.

Libraries of compounds may be presented in solution (e.g., Houghten (1992) Biotechniques 13:412-421), or on beads (Lam (1991) Nature 354:82-84), on chips (Fodor (1993) Nature 364:555-556), bacteria (Ladner U.S. Pat. No. 5,223,409), spores (Ladner USP '409), plasmids (Cull et al. (1992) Proc Natl Acad Sci USA 89:1865-1869) or on phage (Scott and

Smith (1990) Science 249:386-390; Devlin (1990) Science 249:404-406; Cwirla et al. (1990) Proc Natl Acad Sci U.S.A. 87:6378-6382; Felici (1991) J Mol Biol 222:301-310; Ladner above.).

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In one embodiment, an assay is a cell-based assay in which a cell which expresses a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface is contacted with a test compound and the ability of the test compound to bind to a ORFX protein determined. The cell, for example, can of mammalian origin or a yeast cell. Determining the ability of the test compound to bind to the ORFX protein can be accomplished, for example, by coupling the test compound with a radioisotope or enzymatic label such that binding of the test compound to the ORFX protein or biologically active portion thereof can be determined by detecting the labeled compound in a complex. For example, test compounds can be labeled with ¹²⁵I, ³⁵S, ¹⁴C, or ³H, either directly or indirectly, and the radioisotope detected by direct counting of radioemission or by scintillation counting. Alternatively, test compounds can be enzymatically labeled with, for example, horseradish peroxidase, alkaline phosphatase, or luciferase, and the enzymatic label detected by determination of conversion of an appropriate substrate to product. In one embodiment, the assay comprises contacting a cell which expresses a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the test compound to preferentially bind to ORFX or a biologically active portion thereof as compared to the known compound.

In another embodiment, an assay is a cell-based assay comprising contacting a cell expressing a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface with a test compound and determining the ability of the test compound to modulate (e.g., stimulate or inhibit) the activity of the ORFX protein or biologically active portion thereof. Determining the ability of the test compound to modulate the activity of ORFX or a biologically active portion thereof can be accomplished, for example, by determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule. As used herein, a "target molecule" is a molecule with which a ORFX protein binds or interacts in nature, for example, a molecule on the surface of a cell which expresses a ORFX interacting protein, a molecule on the surface of a second cell, a molecule in the extracellular milieu, a molecule

associated with the internal surface of a cell membrane or a cytoplasmic molecule. A ORFX target molecule can be a non-ORFX molecule or a ORFX protein or polypeptide of the present invention. In one embodiment, a ORFX target molecule is a component of a signal transduction pathway that facilitates transduction of an extracellular signal (e.g., a signal generated by binding of a compound to a membrane-bound ORFX molecule) through the cell membrane and into the cell. The target, for example, can be a second intercellular protein that has catalytic activity or a protein that facilitates the association of downstream signaling molecules with ORFX.

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Determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule can be accomplished by one of the methods described above for determining direct binding. In one embodiment, determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule can be accomplished by determining the activity of the target molecule. For example, the activity of the target molecule can be determined by detecting induction of a cellular second messenger of the target (*i.e.* intracellular Ca²⁺, diacylglycerol, IP₃, etc.), detecting catalytic/enzymatic activity of the target an appropriate substrate, detecting the induction of a reporter gene (comprising a ORFX-responsive regulatory element operatively linked to a nucleic acid encoding a detectable marker, *e.g.*, luciferase), or detecting a cellular response, for example, cell survival, cellular differentiation, or cell proliferation.

In yet another embodiment, an assay of the present invention is a cell-free assay comprising contacting a ORFX protein or biologically active portion thereof with a test compound and determining the ability of the test compound to bind to the ORFX protein or biologically active portion thereof. Binding of the test compound to the ORFX protein can be determined either directly or indirectly as described above. In one embodiment, the assay comprises contacting the ORFX protein or biologically active portion thereof with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the test compound to preferentially bind to ORFX or biologically active portion thereof as compared to the known compound.

In another embodiment, an assay is a cell-free assay comprising contacting ORFX protein or biologically active portion thereof with a test compound and determining the ability of the test compound to modulate (e.g., stimulate or inhibit) the activity of the ORFX protein or

biologically active portion thereof. Determining the ability of the test compound to modulate the activity of ORFX can be accomplished, for example, by determining the ability of the ORFX protein to bind to a ORFX target molecule by one of the methods described above for determining direct binding. In an alternative embodiment, determining the ability of the test compound to modulate the activity of ORFX can be accomplished by determining the ability of the ORFX protein further modulate a ORFX target molecule. For example, the catalytic/enzymatic activity of the target molecule on an appropriate substrate can be determined as previously described.

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In yet another embodiment, the cell-free assay comprises contacting the ORFX protein or biologically active portion thereof with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the ORFX protein to preferentially bind to or modulate the activity of a ORFX target molecule.

The cell-free assays of the present invention are amenable to use of both the soluble form or the membrane-bound form of ORFX. In the case of cell-free assays comprising the membrane-bound form of ORFX, it may be desirable to utilize a solubilizing agent such that the membrane-bound form of ORFX is maintained in solution. Examples of such solubilizing agents include non-ionic detergents such as n-octylglucoside, n-dodecylglucoside, n-dodecylmaltoside, octanoyl-N-methylglucamide, decanoyl-N-methylglucamide, Triton® X-100, Triton® X-114, Thesit®, Isotridecypoly(ethylene glycol ether)_n, N-dodecyl--N,N-dimethyl-3-ammonio-1-propane sulfonate, 3-(3-cholamidopropyl)dimethylamminiol-1-propane sulfonate (CHAPSO).

In more than one embodiment of the above assay methods of the present invention, it may be desirable to immobilize either ORFX or its target molecule to facilitate separation of complexed from uncomplexed forms of one or both of the proteins, as well as to accommodate automation of the assay. Binding of a test compound to ORFX, or interaction of ORFX with a target molecule in the presence and absence of a candidate compound, can be accomplished in any vessel suitable for containing the reactants. Examples of such vessels include microtiter plates, test tubes, and micro-centrifuge tubes. In one embodiment, a fusion protein can be provided that adds a domain that allows one or both of the proteins to be bound to a matrix. For

example, GST-ORFX fusion proteins or GST-target fusion proteins can be adsorbed onto glutathione sepharose beads (Sigma Chemical, St. Louis, MO) or glutathione derivatized microtiter plates, that are then combined with the test compound or the test compound and either the non-adsorbed target protein or ORFX protein, and the mixture is incubated under conditions conducive to complex formation (e.g., at physiological conditions for salt and pH). Following incubation, the beads or microtiter plate wells are washed to remove any unbound components, the matrix immobilized in the case of beads, complex determined either directly or indirectly, for example, as described above. Alternatively, the complexes can be dissociated from the matrix, and the level of ORFX binding or activity determined using standard techniques.

Other techniques for immobilizing proteins on matrices can also be used in the screening assays of the invention. For example, either ORFX or its target molecule can be immobilized utilizing conjugation of biotin and streptavidin. Biotinylated ORFX or target molecules can be prepared from biotin-NHS (N-hydroxy-succinimide) using techniques well known in the art (e.g., biotinylation kit, Pierce Chemicals, Rockford, Ill.), and immobilized in the wells of streptavidin-coated 96 well plates (Pierce Chemical). Alternatively, antibodies reactive with ORFX or target molecules, but which do not interfere with binding of the ORFX protein to its target molecule, can be derivatized to the wells of the plate, and unbound target or ORFX trapped in the wells by antibody conjugation. Methods for detecting such complexes, in addition to those described above for the GST-immobilized complexes, include immunodetection of complexes using antibodies reactive with the ORFX or target molecule, as well as enzyme-linked assays that rely on detecting an enzymatic activity associated with the ORFX or target molecule.

In another embodiment, modulators of ORFX expression are identified in a method wherein a cell is contacted with a candidate compound and the expression of ORFX mRNA or protein in the cell is determined. The level of expression of ORFX mRNA or protein in the presence of the candidate compound is compared to the level of expression of ORFX mRNA or protein in the absence of the candidate compound. The candidate compound can then be identified as a modulator of ORFX expression based on this comparison. For example, when expression of ORFX mRNA or protein is greater (statistically significantly greater) in the presence of the candidate compound than in its absence, the candidate compound is identified as a stimulator of ORFX mRNA or protein expression. Alternatively, when expression of ORFX mRNA or protein is less (statistically significantly less) in the presence of the candidate

compound than in its absence, the candidate compound is identified as an inhibitor of ORFX mRNA or protein expression. The level of ORFX mRNA or protein expression in the cells can be determined by methods described herein for detecting ORFX mRNA or protein.

In yet another aspect of the invention, the ORFX proteins can be used as "bait proteins" in a two-hybrid assay or three hybrid assay (see, e.g., U.S. Pat. No. 5,283,317; Zervos et al. (1993) Cell 72:223-232; Madura et al. (1993) J Biol Chem 268:12046-12054; Bartel et al. (1993) Biotechniques 14:920-924; Iwabuchi et al. (1993) Oncogene 8:1693-1696; and Brent WO94/10300), to identify other proteins that bind to or interact with ORFX ("ORFX-binding proteins" or "ORFX-bp") and modulate ORFX activity. Such ORFX-binding proteins are also likely to be involved in the propagation of signals by the ORFX proteins as, for example, upstream or downstream elements of the ORFX pathway.

The two-hybrid system is based on the modular nature of most transcription factors, which consist of separable DNA-binding and activation domains. Briefly, the assay utilizes two different DNA constructs. In one construct, the gene that codes for ORFX is fused to a gene encoding the DNA binding domain of a known transcription factor (e.g., GAL-4). In the other construct, a DNA sequence, from a library of DNA sequences, that encodes an unidentified protein ("prey" or "sample") is fused to a gene that codes for the activation domain of the known transcription factor. If the "bait" and the "prey" proteins are able to interact, in vivo, forming a ORFX-dependent complex, the DNA-binding and activation domains of the transcription factor are brought into close proximity. This proximity allows transcription of a reporter gene (e.g., LacZ) that is operably linked to a transcriptional regulatory site responsive to the transcription factor. Expression of the reporter gene can be detected and cell colonies containing the functional transcription factor can be isolated and used to obtain the cloned gene that encodes the protein which interacts with ORFX.

This invention further pertains to novel agents identified by the above-described screening assays and uses thereof for treatments as described herein.

Detection Assays

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Portions or fragments of the cDNA sequences identified herein (and the corresponding complete gene sequences) can be used in numerous ways as polynucleotide reagents. For example, these sequences can be used to: (i) map their respective genes on a chromosome; and, thus, locate gene regions associated with genetic disease; (ii) identify an individual from a

minute biological sample (tissue typing); and (iii) aid in forensic identification of a biological sample.

The ORFX sequences of the present invention can also be used to identify individuals from minute biological samples. In this technique, an individual's genomic DNA is digested with one or more restriction enzymes, and probed on a Southern blot to yield unique bands for identification. The sequences of the present invention are useful as additional DNA markers for RFLP ("restriction fragment length polymorphisms," described in U.S. Pat. No. 5,272,057).

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Furthermore, the sequences of the present invention can be used to provide an alternative technique that determines the actual base-by-base DNA sequence of selected portions of an individual's genome. Thus, the ORFX sequences described herein can be used to prepare two PCR primers from the 5' and 3' ends of the sequences. These primers can then be used to amplify an individual's DNA and subsequently sequence it.

Panels of corresponding DNA sequences from individuals, prepared in this manner, can provide unique individual identifications, as each individual will have a unique set of such DNA sequences due to allelic differences. The sequences of the present invention can be used to obtain such identification sequences from individuals and from tissue. The ORFX sequences of the invention uniquely represent portions of the human genome. Allelic variation occurs to some degree in the coding regions of these sequences, and to a greater degree in the noncoding regions. It is estimated that allelic variation between individual humans occurs with a frequency of about once per each 500 bases. Much of the allelic variation is due to single nucleotide polymorphisms (SNPs), which include restriction fragment length polymorphisms (RFLPs).

Each of the sequences described herein can, to some degree, be used as a standard against which DNA from an individual can be compared for identification purposes. Because greater numbers of polymorphisms occur in the noncoding regions, fewer sequences are necessary to differentiate individuals. The noncoding sequences of SEQ ID NO:2n-1 (wherein n=1 to 3161), as described above, can comfortably provide positive individual identification with a panel of perhaps 10 to 1,000 primers that each yield a noncoding amplified sequence of 100 bases. If predicted coding sequences are used, a more appropriate number of primers for positive individual identification would be 500-2,000.

Predictive Medicine

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The present invention also pertains to the field of predictive medicine in which diagnostic assays, prognostic assays, pharmacogenomics, and monitoring clinical trials are used for prognostic (predictive) purposes to thereby treat an individual prophylactically. Accordingly, one aspect of the present invention relates to diagnostic assays for determining ORFX protein and/or nucleic acid expression as well as ORFX activity, in the context of a biological sample (e.g., blood, serum, cells, tissue) to thereby determine whether an individual is afflicted with a disease or disorder, or is at risk of developing a disorder, associated with aberrant ORFX expression or activity. The invention also provides for prognostic (or predictive) assays for determining whether an individual is at risk of developing a disorder associated with ORFX protein, nucleic acid expression or activity. For example, mutations in a ORFX gene can be assayed in a biological sample. Such assays can be used for prognostic or predictive purpose to thereby prophylactically treat an individual prior to the onset of a disorder characterized by or associated with ORFX protein, nucleic acid expression or activity.

Another aspect of the invention provides methods for determining ORFX protein, nucleic acid expression or ORFX activity in an individual to thereby select appropriate therapeutic or prophylactic agents for that individual (referred to herein as "pharmacogenomics"). Pharmacogenomics allows for the selection of agents (e.g., drugs) for therapeutic or prophylactic treatment of an individual based on the genotype of the individual (e.g., the genotype of the individual examined to determine the ability of the individual to respond to a particular agent.)

Yet another aspect of the invention pertains to monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX in clinical trials.

Use of Partial ORFX Sequences in Forensic Biology

DNA-based identification techniques can also be used in forensic biology. Forensic biology is a scientific field employing genetic typing of biological evidence found at a crime scene as a means for positively identifying, for example, a perpetrator of a crime. To make such an identification, PCR technology can be used to amplify DNA sequences taken from very small biological samples such as tissues, e.g., hair or skin, or body fluids, e.g., blood, saliva, or semen

found at a crime scene. The amplified sequence can then be compared to a standard, thereby allowing identification of the origin of the biological sample.

The sequences of the present invention can be used to provide polynucleotide reagents, e.g., PCR primers, targeted to specific loci in the human genome, that can enhance the reliability of DNA-based forensic identifications by, for example, providing another "identification marker" (i.e. another DNA sequence that is unique to a particular individual). As mentioned above, actual base sequence information can be used for identification as an accurate alternative to patterns formed by restriction enzyme generated fragments. Sequences targeted to noncoding regions of SEQ ID NOs: __ are particularly appropriate for this use as greater numbers of polymorphisms occur in the noncoding regions, making it easier to differentiate individuals using this technique. Examples of polynucleotide reagents include the ORFX sequences or portions thereof, e.g., fragments derived from the noncoding regions of one or more of SEQ ID NO:2n-1 (where n = 1 to 3161), having a length of at least 20 bases, preferably at least 30 bases.

The ORFX sequences described herein can further be used to provide polynucleotide reagents, e.g., labeled or label-able probes that can be used, for example, in an in situ hybridization technique, to identify a specific tissue, e.g., brain tissue, etc. This can be very useful in cases where a forensic pathologist is presented with a tissue of unknown origin. Panels of such ORFX probes can be used to identify tissue by species and/or by organ type.

In a similar fashion, these reagents, e.g., ORFX primers or probes can be used to screen tissue culture for contamination (i.e. screen for the presence of a mixture of different types of cells in a culture).

Predictive Medicine

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The present invention also pertains to the field of predictive medicine in which diagnostic assays, prognostic assays, pharmacogenomics, and monitoring clinical trials are used for prognostic (predictive) purposes to thereby treat an individual prophylactically. Accordingly, one aspect of the present invention relates to diagnostic assays for determining ORFX protein and/or nucleic acid expression as well as ORFX activity, in the context of a biological sample (e.g., blood, serum, cells, tissue) to thereby determine whether an individual is afflicted with a disease or disorder, or is at risk of developing a disorder, associated with aberrant ORFX expression or activity. The invention also provides for prognostic (or predictive) assays for determining whether an individual is at risk of developing a disorder associated with ORFX protein, nucleic

acid expression or activity. For example, mutations in a ORFX gene can be assayed in a biological sample. Such assays can be used for prognostic or predictive purpose to thereby prophylactically treat an individual prior to the onset of a disorder characterized by or associated with ORFX protein, nucleic acid expression or activity.

Another aspect of the invention provides methods for determining ORFX protein, nucleic acid expression or ORFX activity in an individual to thereby select appropriate therapeutic or prophylactic agents for that individual (referred to herein as "pharmacogenomics"). Pharmacogenomics allows for the selection of agents (e.g., drugs) for therapeutic or prophylactic treatment of an individual based on the genotype of the individual (e.g., the genotype of the individual examined to determine the ability of the individual to respond to a particular agent.)

Yet another aspect of the invention pertains to monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX in clinical trials.

These and other agents are described in further detail in the following sections.

Diagnostic Assays

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Other conditions in which proliferation of cells plays a role include tumors, restenosis, psoriasis, Dupuytren's contracture, diabetic complications, Kaposi's sarcoma and rheumatoid arthritis.

An ORFX polypeptide may be used to identify an interacting polypeptide a sample or tissue. The method comprises contacting the sample or tissue with ORFX, allowing formation of a complex between the ORFX polypeptide and the interacting polypeptide, and detecting the complex, if present.

The proteins of the invention may be used to stimulate production of antibodies specifically binding the proteins. Such antibodies may be used in immunodiagnostic procedures to detect the occurrence of the protein in a sample. The proteins of the invention may be used to stimulate cell growth and cell proliferation in conditions in which such growth would be favorable. An example would be to counteract toxic side effects of chemotherapeutic agents on, for example, hematopoiesis and platelet formation, linings of the gastrointestinal tract, and hair follicles. They may also be used to stimulate new cell growth in neurological disorders including, for example, Alzheimer's disease. Alternatively, antagonistic treatments may be administered in which an antibody specifically binding the ORFX -like proteins of the invention

would abrogate the specific growth-inducing effects of the proteins. Such antibodies may be useful, for example, in the treatment of proliferative disorders including various tumors and benign hyperplasias.

Polynucleotides or oligonucleotides corresponding to any one portion of the ORFX nucleic acids of SEQ ID NO:2n-1 (wherein n=1 to 3161) may be used to detect DNA containing a corresponding ORF gene, or detect the expression of a corresponding ORFX gene, or ORFX-like gene. For example, an ORFX nucleic acid expressed in a particular cell or tissue, as noted in Table 2, can be used to identify the presence of that particular cell type.

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An exemplary method for detecting the presence or absence of ORFX in a biological sample involves obtaining a biological sample from a test subject and contacting the biological sample with a compound or an agent capable of detecting ORFX protein or nucleic acid (e.g., mRNA, genomic DNA) that encodes ORFX protein such that the presence of ORFX is detected in the biological sample. An agent for detecting ORFX mRNA or genomic DNA is a labeled nucleic acid probe capable of hybridizing to ORFX mRNA or genomic DNA. The nucleic acid probe can be, for example, a full-length ORFX nucleic acid, such as the nucleic acid of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a portion thereof, such as an oligonucleotide of at least 15, 30, 50, 100, 250 or 500 nucleotides in length and sufficient to specifically hybridize under stringent conditions to ORFX mRNA or genomic DNA, as described above. Other suitable probes for use in the diagnostic assays of the invention are described herein.

An agent for detecting ORFX protein is an antibody capable of binding to ORFX protein, preferably an antibody with a detectable label. Antibodies can be polyclonal, or more preferably, monoclonal. An intact antibody, or a fragment thereof (e.g., Fab or F(ab')₂) can be used. The term "labeled", with regard to the probe or antibody, is intended to encompass direct labeling of the probe or antibody by coupling (i.e., physically linking) a detectable substance to the probe or antibody, as well as indirect labeling of the probe or antibody by reactivity with another reagent that is directly labeled. Examples of indirect labeling include detection of a primary antibody using a fluorescently labeled secondary antibody and end-labeling of a DNA probe with biotin such that it can be detected with fluorescently labeled streptavidin. The term "biological sample" is intended to include tissues, cells and biological fluids isolated from a subject, as well as tissues, cells and fluids present within a subject. That is, the detection method of the invention can be used to detect ORFX mRNA, protein, or genomic DNA in a biological sample in vitro as

well as *in vivo*. For example, *in vitro* techniques for detection of ORFX mRNA include Northern hybridizations and *in situ* hybridizations. *In vitro* techniques for detection of ORFX protein include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations and immunofluorescence. *In vitro* techniques for detection of ORFX genomic DNA include Southern hybridizations. Furthermore, *in vivo* techniques for detection of ORFX protein include introducing into a subject a labeled anti-ORFX antibody. For example, the antibody can be labeled with a radioactive marker whose presence and location in a subject can be detected by standard imaging techniques.

In one embodiment, the biological sample contains protein molecules from the test subject. Alternatively, the biological sample can contain mRNA molecules from the test subject or genomic DNA molecules from the test subject. A preferred biological sample is a peripheral blood leukocyte sample isolated by conventional means from a subject.

In another embodiment, the methods further involve obtaining a control biological sample from a control subject, contacting the control sample with a compound or agent capable of detecting ORFX protein, mRNA, or genomic DNA, such that the presence of ORFX protein, mRNA or genomic DNA is detected in the biological sample, and comparing the presence of ORFX protein, mRNA or genomic DNA in the control sample with the presence of ORFX protein, mRNA or genomic DNA in the test sample.

The invention also encompasses kits for detecting the presence of ORFX in a biological sample. For example, the kit can comprise: a labeled compound or agent capable of detecting ORFX protein or mRNA in a biological sample; means for determining the amount of ORFX in the sample; and means for comparing the amount of ORFX in the sample with a standard. The compound or agent can be packaged in a suitable container. The kit can further comprise instructions for using the kit to detect ORFX protein or nucleic acid.

Prognostic Assays

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The diagnostic methods described herein can furthermore be utilized to identify subjects having or at risk of developing a disease or disorder associated with aberrant ORFX expression or activity. For example, the assays described herein, such as the preceding diagnostic assays or the following assays, can be utilized to identify a subject having or at risk of developing a disorder associated with ORFX protein, nucleic acid expression or activity in, e.g., proliferative or differentiative disorders such as hyperplasias, tumors, restenosis, psoriasis, Dupuytren's

contracture, diabetic complications, or rheumatoid arthritis, etc.; and glia-associated disorders such as cerebral lesions, diabetic neuropathies, cerebral edema, senile dementia, Alzheimer's disease, etc. Alternatively, the prognostic assays can be utilized to identify a subject having or at risk for developing a disease or disorder. Thus, the present invention provides a method for identifying a disease or disorder associated with aberrant ORFX expression or activity in which a test sample is obtained from a subject and ORFX protein or nucleic acid (e.g., mRNA, genomic DNA) is detected, wherein the presence of ORFX protein or nucleic acid is diagnostic for a subject having or at risk of developing a disease or disorder associated with aberrant ORFX expression or activity. As used herein, a "test sample" refers to a biological sample obtained from a subject of interest. For example, a test sample can be a biological fluid (e.g., serum), cell sample, or tissue.

Furthermore, the prognostic assays described herein can be used to determine whether a subject can be administered an agent (e.g., an agonist, antagonist, peptidomimetic, protein, peptide, nucleic acid, small molecule, or other drug candidate) to treat a disease or disorder associated with aberrant ORFX expression or activity. For example, such methods can be used to determine whether a subject can be effectively treated with an agent for a disorder, such as a proliferative disorder, differentiative disorder, glia-associated disorders, etc. Thus, the present invention provides methods for determining whether a subject can be effectively treated with an agent for a disorder associated with aberrant ORFX expression or activity in which a test sample is obtained and ORFX protein or nucleic acid is detected (e.g., wherein the presence of ORFX protein or nucleic acid is diagnostic for a subject that can be administered the agent to treat a disorder associated with aberrant ORFX expression or activity.)

The methods of the invention can also be used to detect genetic lesions in a ORFX gene, thereby determining if a subject with the lesioned gene is at risk for, or suffers from, a proliferative disorder, differentiative disorder, glia-associated disorder, etc. In various embodiments, the methods include detecting, in a sample of cells from the subject, the presence or absence of a genetic lesion characterized by at least one of an alteration affecting the integrity of a gene encoding a ORFX-protein, or the mis-expression of the ORFX gene. For example, such genetic lesions can be detected by ascertaining the existence of at least one of (1) a deletion of one or more nucleotides from a ORFX gene; (2) an addition of one or more nucleotides to a ORFX gene; (3) a substitution of one or more nucleotides of a ORFX gene, (4) a chromosomal

rearrangement of a ORFX gene; (5) an alteration in the level of a messenger RNA transcript of a ORFX gene, (6) aberrant modification of a ORFX gene, such as of the methylation pattern of the genomic DNA, (7) the presence of a non-wild type splicing pattern of a messenger RNA transcript of a ORFX gene, (8) a non-wild type level of a ORFX-protein, (9) allelic loss of a ORFX gene, and (10) inappropriate post-translational modification of a ORFX-protein. As described herein, there are a large number of assay techniques known in the art which can be used for detecting lesions in a ORFX gene. A preferred biological sample is a peripheral blood leukocyte sample isolated by conventional means from a subject. However, any biological sample containing nucleated cells may be used, including, for example, buccal mucosal cells.

In certain embodiments, detection of the lesion involves the use of a probe/primer in a polymerase chain reaction (PCR) (see, e.g., U.S. Pat. Nos. 4,683,195 and 4,683,202), such as anchor PCR or RACE PCR, or, alternatively, in a ligation chain reaction (LCR) (see, e.g., Landegran et al. (1988) Science 241:1077-1080; and Nakazawa et al. (1994) PNAS 91:360-364), the latter of which can be particularly useful for detecting point mutations in the ORFX-gene (see Abravaya et al. (1995) Nucl Acids Res 23:675-682). This method can include the steps of collecting a sample of cells from a patient, isolating nucleic acid (e.g., genomic, mRNA or both) from the cells of the sample, contacting the nucleic acid sample with one or more primers that specifically hybridize to a ORFX gene under conditions such that hybridization and amplification of the ORFX gene (if present) occurs, and detecting the presence or absence of an amplification product, or detecting the size of the amplification product and comparing the length to a control sample. It is anticipated that PCR and/or LCR may be desirable to use as a preliminary amplification step in conjunction with any of the techniques used for detecting mutations described herein.

Alternative amplification methods include: self sustained sequence replication (Guatelli et al., 1990, Proc Natl Acad Sci USA 87:1874-1878), transcriptional amplification system (Kwoh, et al., 1989, Proc Natl Acad Sci USA 86:1173-1177), Q-Beta Replicase (Lizardi et al, 1988, BioTechnology 6:1197), or any other nucleic acid amplification method, followed by the detection of the amplified molecules using techniques well known to those of skill in the art. These detection schemes are especially useful for the detection of nucleic acid molecules if such molecules are present in very low numbers.

In an alternative embodiment, mutations in a ORFX gene from a sample cell can be identified by alterations in restriction enzyme cleavage patterns. For example, sample and control DNA is isolated, amplified (optionally), digested with one or more restriction endonucleases, and fragment length sizes are determined by gel electrophoresis and compared. Differences in fragment length sizes between sample and control DNA indicates mutations in the sample DNA. Moreover, the use of sequence specific ribozymes (see, for example, U.S. Pat. No. 5,493,531) can be used to score for the presence of specific mutations by development or loss of a ribozyme cleavage site.

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In other embodiments, genetic mutations in ORFX can be identified by hybridizing a sample and control nucleic acids, e.g., DNA or RNA, to high density arrays containing hundreds or thousands of oligonucleotides probes (Cronin et al. (1996) Human Mutation 7: 244-255; Kozal et al. (1996) Nature Medicine 2: 753-759). For example, genetic mutations in ORFX can be identified in two dimensional arrays containing light-generated DNA probes as described in Cronin et al. above. Briefly, a first hybridization array of probes can be used to scan through long stretches of DNA in a sample and control to identify base changes between the sequences by making linear arrays of sequential overlapping probes. This step allows the identification of point mutations. This step is followed by a second hybridization array that allows the characterization of specific mutations by using smaller, specialized probe arrays complementary to all variants or mutations detected. Each mutation array is composed of parallel probe sets, one complementary to the wild-type gene and the other complementary to the mutant gene.

In yet another embodiment, any of a variety of sequencing reactions known in the art can be used to directly sequence the ORFX gene and detect mutations by comparing the sequence of the sample ORFX with the corresponding wild-type (control) sequence. Examples of sequencing reactions include those based on techniques developed by Maxim and Gilbert (1977) *PNAS* 74:560 or Sanger (1977) *PNAS* 74:5463. It is also contemplated that any of a variety of automated sequencing procedures can be utilized when performing the diagnostic assays (Naeve et al., (1995) Biotechniques 19:448), including sequencing by mass spectrometry (see, e.g., PCT International Publ. No. WO 94/16101; Cohen et al. (1996) Adv Chromatogr 36:127-162; and Griffin et al. (1993) Appl Biochem Biotechnol 38:147-159).

Other methods for detecting mutations in the ORFX gene include methods in which protection from cleavage agents is used to detect mismatched bases in RNA/RNA or RNA/DNA

mismatch cleavage" starts by providing heteroduplexes of formed by hybridizing (labeled) RNA or DNA containing the wild-type ORFX sequence with potentially mutant RNA or DNA obtained from a tissue sample. The double-stranded duplexes are treated with an agent that cleaves single-stranded regions of the duplex such as which will exist due to basepair mismatches between the control and sample strands. For instance, RNA/DNA duplexes can be treated with RNase and DNA/DNA hybrids treated with S1 nuclease to enzymatically digesting the mismatched regions. In other embodiments, either DNA/DNA or RNA/DNA duplexes can be treated with hydroxylamine or osmium tetroxide and with piperidine in order to digest mismatched regions. After digestion of the mismatched regions, the resulting material is then separated by size on denaturing polyacrylamide gels to determine the site of mutation. See, for example, Cotton et al (1988) Proc Natl Acad Sci USA 85:4397; Saleeba et al (1992) Methods Enzymol 217:286-295. In an embodiment, the control DNA or RNA can be labeled for detection.

In still another embodiment, the mismatch cleavage reaction employs one or more proteins that recognize mismatched base pairs in double-stranded DNA (so called "DNA mismatch repair" enzymes) in defined systems for detecting and mapping point mutations in ORFX cDNAs obtained from samples of cells. For example, the mutY enzyme of *E. coli* cleaves A at G/A mismatches and the thymidine DNA glycosylase from HeLa cells cleaves T at G/T mismatches (Hsu et al. (1994) Carcinogenesis 15:1657-1662). According to an exemplary embodiment, a probe based on a ORFX sequence, e.g., a wild-type ORFX sequence, is hybridized to a cDNA or other DNA product from a test cell(s). The duplex is treated with a DNA mismatch repair enzyme, and the cleavage products, if any, can be detected from electrophoresis protocols or the like. See, for example, U.S. Pat. No. 5,459,039.

In other embodiments, alterations in electrophoretic mobility will be used to identify mutations in ORFX genes. For example, single strand conformation polymorphism (SSCP) may be used to detect differences in electrophoretic mobility between mutant and wild type nucleic acids (Orita et al. (1989) Proc Natl Acad Sci USA: 86:2766, see also Cotton (1993) Mutat Res 285:125-144; Hayashi (1992) Genet Anal Tech Appl 9:73-79). Single-stranded DNA fragments of sample and control ORFX nucleic acids will be denatured and allowed to renature. The secondary structure of single-stranded nucleic acids varies according to sequence, the resulting

alteration in electrophoretic mobility enables the detection of even a single base change. The DNA fragments may be labeled or detected with labeled probes. The sensitivity of the assay may be enhanced by using RNA, rather than DNA, in which the secondary structure is more sensitive to a change in sequence. In one embodiment, the subject method utilizes heteroduplex analysis to separate double stranded heteroduplex molecules on the basis of changes in electrophoretic mobility. See, e.g., Keen et al. (1991) Trends Genet 7:5.

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In yet another embodiment the movement of mutant or wild-type fragments in polyacrylamide gels containing a gradient of denaturant is assayed using denaturing gradient gel electrophoresis (DGGE). See, e.g., Myers et al (1985) Nature 313:495. When DGGE is used as the method of analysis, DNA will be modified to insure that it does not completely denature, for example by adding a GC clamp of approximately 40 bp of high-melting GC-rich DNA by PCR. In a further embodiment, a temperature gradient is used in place of a denaturing gradient to identify differences in the mobility of control and sample DNA. See, e.g., Rosenbaum and Reissner (1987) Biophys Chem 265:12753.

Examples of other techniques for detecting point mutations include, but are not limited to, selective oligonucleotide hybridization, selective amplification, or selective primer extension. For example, oligonucleotide primers may be prepared in which the known mutation is placed centrally and then hybridized to target DNA under conditions that permit hybridization only if a perfect match is found. See, e.g., Saiki et al. (1986) Nature 324:163); Saiki et al. (1989) Proc Natl Acad. Sci USA 86:6230. Such allele specific oligonucleotides are hybridized to PCR amplified target DNA or a number of different mutations when the oligonucleotides are attached to the hybridizing membrane and hybridized with labeled target DNA.

Alternatively, allele specific amplification technology that depends on selective PCR amplification may be used in conjunction with the instant invention. Oligonucleotides used as primers for specific amplification may carry the mutation of interest in the center of the molecule (so that amplification depends on differential hybridization) (Gibbs et al. (1989) Nucleic Acids Res 17:2437-2448) or at the extreme 3' end of one primer where, under appropriate conditions, mismatch can prevent, or reduce polymerase extension (Prossner (1993) Tibtech 11:238). In addition it may be desirable to introduce a novel restriction site in the region of the mutation to create cleavage-based detection. See, e.g., Gasparini et al (1992) Mol Cell Probes 6:1. It is anticipated that in certain embodiments amplification may also be performed using Taq ligase

for amplification. See, e.g., Barany (1991) Proc Natl Acad Sci USA 88:189. In such cases, ligation will occur only if there is a perfect match at the 3' end of the 5' sequence, making it possible to detect the presence of a known mutation at a specific site by looking for the presence or absence of amplification.

The methods described herein may be performed, for example, by utilizing pre-packaged diagnostic kits comprising at least one probe nucleic acid or antibody reagent described herein, which may be conveniently used, e.g., in clinical settings to diagnose patients exhibiting symptoms or family history of a disease or illness involving a ORFX gene.

Furthermore, any cell type or tissue, preferably peripheral blood leukocytes, in which ORFX is expressed may be utilized in the prognostic assays described herein. However, any biological sample containing nucleated cells may be used, including, for example, buccal mucosal cells.

Pharmacogenomics

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Agents, or modulators that have a stimulatory or inhibitory effect on ORFX activity (e.g., ORFX gene expression), as identified by a screening assay described herein can be administered to individuals to treat (prophylactically or therapeutically) disorders (e.g., neurological, cancerrelated or gestational disorders) associated with aberrant ORFX activity. In conjunction with such treatment, the pharmacogenomics (i.e., the study of the relationship between an individual's genotype and that individual's response to a foreign compound or drug) of the individual may be considered. Differences in metabolism of therapeutics can lead to severe toxicity or therapeutic failure by altering the relation between dose and blood concentration of the pharmacologically active drug. Thus, the pharmacogenomics of the individual permits the selection of effective agents (e.g., drugs) for prophylactic or therapeutic treatments based on a consideration of the individual's genotype. Such pharmacogenomics can further be used to determine appropriate dosages and therapeutic regimens. Accordingly, the activity of ORFX protein, expression of ORFX nucleic acid, or mutation content of ORFX genes in an individual can be determined to thereby select appropriate agent(s) for therapeutic or prophylactic treatment of the individual.

Pharmacogenomics deals with clinically significant hereditary variations in the response to drugs due to altered drug disposition and abnormal action in affected persons. See e.g., Eichelbaum, 1996, Clin Exp Pharmacol Physiol, 23:983-985 and Linder, 1997, Clin Chem, 43:254-266. In general, two types of pharmacogenetic conditions can be differentiated. Genetic

conditions transmitted as a single factor altering the way drugs act on the body (altered drug action) or genetic conditions transmitted as single factors altering the way the body acts on drugs (altered drug metabolism). These pharmacogenetic conditions can occur either as rare defects or as polymorphisms. For example, glucose-6-phosphate dehydrogenase (G6PD) deficiency is a common inherited enzymopathy in which the main clinical complication is haemolysis after ingestion of oxidant drugs (anti-malarials, sulfonamides, analgesics, nitrofurans) and consumption of fava beans.

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As an illustrative embodiment, the activity of drug metabolizing enzymes is a major determinant of both the intensity and duration of drug action. The discovery of genetic polymorphisms of drug metabolizing enzymes (e.g., N-acetyltransferase 2 (NAT 2) and cytochrome P450 enzymes CYP2D6 and CYP2C19) has provided an explanation as to why some patients do not obtain the expected drug effects or show exaggerated drug response and serious toxicity after taking the standard and safe dose of a drug. These polymorphisms are expressed in two phenotypes in the population, the extensive metabolizer (EM) and poor metabolizer (PM). The prevalence of PM is different among different populations. For example, the gene coding for CYP2D6 is highly polymorphic and several mutations have been identified in PM, which all lead to the absence of functional CYP2D6. Poor metabolizers of CYP2D6 and CYP2C19 quite frequently experience exaggerated drug response and side effects when they receive standard doses. If a metabolite is the active therapeutic moiety, PM show no therapeutic response, as demonstrated for the analgesic effect of codeine mediated by its CYP2D6-formed metabolite morphine. The other extreme are the so called ultra-rapid metabolizers who do not respond to standard doses. Recently, the molecular basis of ultra-rapid metabolism has been identified to be due to CYP2D6 gene amplification.

Thus, the activity of ORFX protein, expression of ORFX nucleic acid, or mutation content of ORFX genes in an individual can be determined to thereby select appropriate agent(s) for therapeutic or prophylactic treatment of the individual. In addition, pharmacogenetic studies can be used to apply genotyping of polymorphic alleles encoding drug-metabolizing enzymes to the identification of an individual's drug responsiveness phenotype. This knowledge, when applied to dosing or drug selection, can avoid adverse reactions or therapeutic failure and thus enhance therapeutic or prophylactic efficiency when treating a subject with a ORFX modulator, such as a modulator identified by one of the exemplary screening assays described herein.

Monitoring Clinical Efficacy

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Monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX (e.g., the ability to modulate aberrant cell proliferation and/or differentiation) can be applied in basic drug screening and in clinical trials. For example, the effectiveness of an agent determined by a screening assay as described herein to increase ORFX gene expression, protein levels, or upregulate ORFX activity, can be monitored in clinical trials of subjects exhibiting decreased ORFX gene expression, protein levels, or downregulated ORFX activity.

Alternatively, the effectiveness of an agent determined by a screening assay to decrease ORFX gene expression, protein levels, or downregulate ORFX activity, can be monitored in clinical trials of subjects exhibiting increased ORFX gene expression, protein levels, or upregulated ORFX activity. In such clinical trials, the expression or activity of ORFX and, preferably, other genes that have been implicated in, for example, a proliferative or neurological disorder, can be used as a "read out" or marker of the responsiveness of a particular cell.

For example, genes, including ORFX, that are modulated in cells by treatment with an agent (e.g., compound, drug or small molecule) that modulates ORFX activity (e.g., identified in a screening assay as described herein) can be identified. Thus, to study the effect of agents on cellular proliferation disorders, for example, in a clinical trial, cells can be isolated and RNA prepared and analyzed for the levels of expression of ORFX and other genes implicated in the disorder. The levels of gene expression (i.e., a gene expression pattern) can be quantified by Northern blot analysis or RT-PCR, as described herein, or alternatively by measuring the amount of protein produced, by one of the methods as described herein, or by measuring the levels of activity of ORFX or other genes. In this way, the gene expression pattern can serve as a marker, indicative of the physiological response of the cells to the agent. Accordingly, this response state may be determined before, and at various points during, treatment of the individual with the agent.

In one embodiment, the invention provides a method for monitoring the effectiveness of treatment of a subject with an agent (e.g., an agonist, antagonist, protein, peptide, nucleic acid, peptidomimetic, small molecule, or other drug candidate identified by the screening assays described herein) comprising the steps of (i) obtaining a pre-administration sample from a subject prior to administration of the agent; (ii) detecting the level of expression of a ORFX protein, mRNA, or genomic DNA in the preadministration sample; (iii) obtaining one or more

post-administration samples from the subject; (*iv*) detecting the level of expression or activity of the ORFX protein, mRNA, or genomic DNA in the post-administration samples; (*v*) comparing the level of expression or activity of the ORFX protein, mRNA, or genomic DNA in the pre-administration sample with the ORFX protein, mRNA, or genomic DNA in the post administration sample or samples; and (*vi*) altering the administration of the agent to the subject accordingly. For example, increased administration of the agent may be desirable to increase the expression or activity of ORFX to higher levels than detected, *i.e.*, to increase the effectiveness of the agent. Alternatively, decreased administration of the agent may be desirable to decrease expression or activity of ORFX to lower levels than detected, *i.e.*, to decrease the effectiveness of the agent.

Methods of Treatment

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The present invention provides for both prophylactic and therapeutic methods of treating a subject at risk of (or susceptible to) a disorder or having a disorder associated with aberrant ORFX expression or activity.

Diseases and disorders that are characterized by increased (relative to a subject not suffering from the disease or disorder) levels or biological activity may be treated with Therapeutics that antagonize (*i.e.*, reduce or inhibit) activity. Therapeutics that antagonize activity may be administered in a therapeutic or prophylactic manner. Therapeutics that may be utilized include, but are not limited to, (*i*) a ORFX polypeptide, or analogs, derivatives, fragments or homologs thereof; (*ii*) antibodies to a ORFX peptide; (*iii*) nucleic acids encoding a ORFX peptide; (*iv*) administration of antisense nucleic acid and nucleic acids that are "dysfunctional" (*i.e.*, due to a heterologous insertion within the coding sequences of coding sequences to a ORFX peptide) that are utilized to "knockout" endogenous function of a ORFX peptide by homologous recombination (see, *e.g.*, Capecchi, 1989, *Science* 244: 1288-1292); or (*v*) modulators (*i.e.*, inhibitors, agonists and antagonists, including additional peptide mimetic of the invention or antibodies specific to a peptide of the invention) that alter the interaction between a ORFX peptide and its binding partner.

Diseases and disorders that are characterized by decreased (relative to a subject not suffering from the disease or disorder) levels or biological activity may be treated with Therapeutics that increase (i.e., are agonists to) activity. Therapeutics that upregulate activity may be administered in a therapeutic or prophylactic manner. Therapeutics that may be utilized

include, but are not limited to, a ORFX peptide, or analogs, derivatives, fragments or homologs thereof; or an agonist that increases bioavailability.

Increased or decreased levels can be readily detected by quantifying peptide and/or RNA, by obtaining a patient tissue sample (e.g., from biopsy tissue) and assaying it in vitro for RNA or peptide levels, structure and/or activity of the expressed peptides (or mRNAs of a ORFX peptide). Methods that are well-known within the art include, but are not limited to, immunoassays (e.g., by Western blot analysis, immunoprecipitation followed by sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis, immunocytochemistry, etc.) and/or hybridization assays to detect expression of mRNAs (e.g., Northern assays, dot blots, in situ hybridization, etc.).

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In one aspect, the invention provides a method for preventing, in a subject, a disease or condition associated with an aberrant ORFX expression or activity, by administering to the subject an agent that modulates ORFX expression or at least one ORFX activity. Subjects at risk for a disease that is caused or contributed to by aberrant ORFX expression or activity can be identified by, for example, any or a combination of diagnostic or prognostic assays as described herein. Administration of a prophylactic agent can occur prior to the manifestation of symptoms characteristic of the ORFX aberrancy, such that a disease or disorder is prevented or, alternatively, delayed in its progression. Depending on the type of ORFX aberrancy, for example, a ORFX agonist or ORFX antagonist agent can be used for treating the subject. The appropriate agent can be determined based on screening assays described herein.

Another aspect of the invention pertains to methods of modulating ORFX expression or activity for therapeutic purposes. The modulatory method of the invention involves contacting a cell with an agent that modulates one or more of the activities of ORFX protein activity associated with the cell. An agent that modulates ORFX protein activity can be an agent as described herein, such as a nucleic acid or a protein, a naturally-occurring cognate ligand of a ORFX protein, a peptide, a ORFX peptidomimetic, or other small molecule. In one embodiment, the agent stimulates one or more ORFX protein activity. Examples of such stimulatory agents include active ORFX protein and a nucleic acid molecule encoding ORFX that has been introduced into the cell. In another embodiment, the agent inhibits one or more ORFX protein activity. Examples of such inhibitory agents include antisense ORFX nucleic acid molecules and anti-ORFX antibodies. These modulatory methods can be performed *in vitro*

(e.g., by culturing the cell with the agent) or, alternatively, in vivo (e.g., by administering the agent to a subject). As such, the present invention provides methods of treating an individual afflicted with a disease or disorder characterized by aberrant expression or activity of a ORFX protein or nucleic acid molecule. In one embodiment, the method involves administering an agent (e.g., an agent identified by a screening assay described herein), or combination of agents that modulates (e.g., upregulates or downregulates) ORFX expression or activity. In another embodiment, the method involves administering a ORFX protein or nucleic acid molecule as therapy to compensate for reduced or aberrant ORFX expression or activity.

Determination of the Biological Effect of a Therapeutic

In various embodiments of the present invention, suitable *in vitro* or *in vivo* assays are utilized to determine the effect of a specific Therapeutic and whether its administration is indicated for treatment of the affected tissue.

In various specific embodiments, *in vitro* assays may be performed with representative cells of the type(s) involved in the patient's disorder, to determine if a given Therapeutic exerts the desired effect upon the cell type(s). Compounds for use in therapy may be tested in suitable animal model systems including, but not limited to rats, mice, chicken, cows, monkeys, rabbits, and the like, prior to testing in human subjects. Similarly, for *in vivo* testing, any of the animal model system known in the art may be used prior to administration to human subjects.

Malignancies

Some ORFX polypeptides are expressed in cancerous cells (see, e.g., Tables 1 and 2). Accordingly, the corresponding ORF protein is involved in the regulation of cell proliferation. Accordingly, Therapeutics of the present invention may be useful in the therapeutic or prophylactic treatment of diseases or disorders that are associated with cell hyperproliferation and/or loss of control of cell proliferation (e.g., cancers, malignancies and tumors). For a review of such hyperproliferation disorders, see e.g., Fishman, et al., 1985. MEDICINE, 2nd ed., J.B. Lippincott Co., Philadelphia, PA.

Therapeutics of the present invention may be assayed by any method known within the art for efficacy in treating or preventing malignancies and related disorders. Such assays include,

but are not limited to, *in vitro* assays utilizing transformed cells or cells derived from the patient's tumor, as well as *in vivo* assays using animal models of cancer or malignancies. Potentially effective Therapeutics are those that, for example, inhibit the proliferation of tumor-derived or transformed cells in culture or cause a regression of tumors in animal models, in comparison to the controls.

In the practice of the present invention, once a malignancy or cancer has been shown to be amenable to treatment by modulating (i.e., inhibiting, antagonizing or agonizing) activity, that cancer or malignancy may subsequently be treated or prevented by the administration of a Therapeutic that serves to modulate protein function.

Premalignant conditions

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The Therapeutics of the present invention that are effective in the therapeutic or prophylactic treatment of cancer or malignancies may also be administered for the treatment of pre-malignant conditions and/or to prevent the progression of a pre-malignancy to a neoplastic or malignant state. Such prophylactic or therapeutic use is indicated in conditions known or suspected of preceding progression to neoplasia or cancer, in particular, where non-neoplastic cell growth consisting of hyperplasia, metaplasia or, most particularly, dysplasia has occurred. For a review of such abnormal cell growth see *e.g.*, Robbins & Angell, 1976. Basic Pathology, 2nd ed., W.B. Saunders Co., Philadelphia, PA.

Hyperplasia is a form of controlled cell proliferation involving an increase in cell number in a tissue or organ, without significant alteration in its structure or function. For example, it has been demonstrated that endometrial hyperplasia often precedes endometrial cancer. Metaplasia is a form of controlled cell growth in which one type of mature or fully differentiated cell substitutes for another type of mature cell. Metaplasia may occur in epithelial or connective tissue cells. Dysplasia is generally considered a precursor of cancer, and is found mainly in the epithelia. Dysplasia is the most disorderly form of non-neoplastic cell growth, and involves a loss in individual cell uniformity and in the architectural orientation of cells. Dysplasia characteristically occurs where there exists chronic irritation or inflammation, and is often found in the cervix, respiratory passages, oral cavity, and gall bladder.

Alternatively, or in addition to the presence of abnormal cell growth characterized as hyperplasia, metaplasia, or dysplasia, the presence of one or more characteristics of a transformed or malignant phenotype displayed either *in vivo* or *in vitro* within a cell sample derived from a patient, is indicative of the desirability of prophylactic/therapeutic administration of a Therapeutic that possesses the ability to modulate activity of An aforementioned protein. Characteristics of a transformed phenotype include, but are not limited to: (*i*) morphological changes; (*ii*) looser substratum attachment; (*iii*) loss of cell-to-cell contact inhibition; (*iv*) loss of anchorage dependence; (*v*) protease release; (*vi*) increased sugar transport; (*vii*) decreased serum requirement; (*viii*) expression of fetal antigens, (*ix*) disappearance of the 250 kDal cell-surface protein, and the like. See *e.g.*, Richards, *et al.*, 1986. MOLECULAR PATHOLOGY, W.B. Saunders Co., Philadelphia, PA.

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In a specific embodiment of the present invention, a patient that exhibits one or more of the following predisposing factors for malignancy is treated by administration of an effective amount of a Therapeutic: (i) a chromosomal translocation associated with a malignancy (e.g., the Philadelphia chromosome (bcr/abl) for chronic myelogenous leukemia and t(14;18) for follicular lymphoma, etc.); (ii) familial polyposis or Gardner's syndrome (possible forerunners of colon cancer); (iii) monoclonal gammopathy of undetermined significance (a possible precursor of multiple myeloma) and (iv) a first degree kinship with persons having a cancer or pre-cancerous disease showing a Mendelian (genetic) inheritance pattern (e.g., familial polyposis of the colon, Gardner's syndrome, hereditary exostosis, polyendocrine adenomatosis, Peutz-Jeghers syndrome, neurofibromatosis of Von Recklinghausen, medullary thyroid carcinoma with amyloid production and pheochromocytoma, retinoblastoma, carotid body tumor, cutaneous melanocarcinoma, intraocular melanocarcinoma, xeroderma pigmentosum, ataxia telangiectasia, Chediak-Higashi syndrome, albinism, Fanconi's aplastic anemia and Bloom's syndrome).

In another embodiment, a Therapeutic of the present invention is administered to a human patient to prevent the progression to breast, colon, lung, pancreatic, or uterine cancer, or melanoma or sarcoma.

Hyperproliferative and dysproliferative disorders

In one embodiment of the present invention, a Therapeutic is administered in the therapeutic or prophylactic treatment of hyperproliferative or benign dysproliferative disorders. The efficacy in treating or preventing hyperproliferative diseases or disorders of a Therapeutic of the present invention may be assayed by any method known within the art. Such assays include in vitro cell proliferation assays, in vitro or in vivo assays using animal models of hyperproliferative diseases or disorders, or the like. Potentially effective Therapeutics may, for example, promote cell proliferation in culture or cause growth or cell proliferation in animal models in comparison to controls.

Specific embodiments of the present invention are directed to the treatment or prevention of cirrhosis of the liver (a condition in which scarring has overtaken normal liver regeneration processes); treatment of keloid (hypertrophic scar) formation causing disfiguring of the skin in which the scarring process interferes with normal renewal; psoriasis (a common skin condition characterized by excessive proliferation of the skin and delay in proper cell fate determination); benign tumors; fibrocystic conditions and tissue hypertrophy (e.g., benign prostatic hypertrophy).

Neurodegenerative disorders

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Some ORFX proteins are found in cell types have been implicated in the deregulation of cellular maturation and apoptosis, which are both characteristic of neurodegenerative disease. Accordingly, Therapeutics of the invention, particularly but not limited to those that modulate (or supply) activity of an aforementioned protein, may be effective in treating or preventing neurodegenerative disease. Therapeutics of the present invention that modulate the activity of an aforementioned protein involved in neurodegenerative disorders can be assayed by any method known in the art for efficacy in treating or preventing such neurodegenerative diseases and disorders. Such assays include *in vitro* assays for regulated cell maturation or inhibition of apoptosis or *in vivo* assays using animal models of neurodegenerative diseases or disorders, or any of the assays described below. Potentially effective Therapeutics, for example but not by way of limitation, promote regulated cell maturation and prevent cell apoptosis in culture, or reduce neurodegeneration in animal models in comparison to controls.

Once a neurodegenerative disease or disorder has been shown to be amenable to treatment by modulation activity, that neurodegenerative disease or disorder can be treated or prevented by administration of a Therapeutic that modulates activity. Such diseases include all degenerative disorders involved with aging, especially osteoarthritis and neurodegenerative disorders.

Disorders related to organ transplantation

Some ORFX can be associated with disorders related to organ transplantation, in particular but not limited to organ rejection. Therapeutics of the invention, particularly those that modulate (or supply) activity, may be effective in treating or preventing diseases or disorders related to organ transplantation. Therapeutics of the invention (particularly Therapeutics that modulate the levels or activity of an aforementioned protein) can be assayed by any method known in the art for efficacy in treating or preventing such diseases and disorders related to organ transplantation. Such assays include *in vitro* assays for using cell culture models as described below, or *in vivo* assays using animal models of diseases and disorders related to organ transplantation, see *e.g.*, below. Potentially effective Therapeutics, for example but not by way of limitation, reduce immune rejection responses in animal models in comparison to controls.

Accordingly, once diseases and disorders related to organ transplantation are shown to be amenable to treatment by modulation of activity, such diseases or disorders can be treated or prevented by administration of a Therapeutic that modulates activity.

20 Cardiovascular Disease

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GENX has been implicated in cardiovascular disorders, including in atherosclerotic plaque formation. Diseases such as cardiovascular disease, including cerebral thrombosis or hemorrhage, ischemic heart or renal disease, peripheral vascular disease, or thrombosis of other major vessel, and other diseases, including diabetes mellitus, hypertension, hypothyroidism, cholesterol ester storage disease, systemic lupus erythematosus, homocysteinemia, and familial protein or lipid processing diseases, and the like, are either directly or indirectly associated with atherosclerosis. Accordingly, Therapeutics of the invention, particularly those that modulate (or supply) activity or formation may be effective in treating or preventing

atherosclerosis-associated diseases or disorders. Therapeutics of the invention (particularly Therapeutics that modulate the levels or activity) can be assayed by any method known in the art, including those described below, for efficacy in treating or preventing such diseases and disorders.

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A vast array of animal and cell culture models exist for processes involved in atherosclerosis. A limited and non-exclusive list of animal models includes knockout mice for premature atherosclerosis (Kurabayashi and Yazaki, 1996, Int. Angiol. 15: 187-194), transgenic mouse models of atherosclerosis (Kappel et al., 1994, FASEB J. 8: 583-592), antisense oligonucleotide treatment of animal models (Callow, 1995, Curr. Opin. Cardiol. 10: 569-576), transgenic rabbit models for atherosclerosis (Taylor, 1997, Ann. N.Y. Acad. Sci 811: 146-152), hypercholesterolemic animal models (Rosenfeld, 1996, Diabetes Res. Clin. Pract. 30 Suppl.: 1-11), hyperlipidemic mice (Paigen et al., 1994, Curr. Opin. Lipidol. 5: 258-264), and inhibition of lipoxygenase in animals (Sigal et al., 1994, Ann. N.Y. Acad. Sci. 714: 211-224). In addition, in vitro cell models include but are not limited to monocytes exposed to low density lipoprotein (Frostegard et al., 1996, Atherosclerosis 121: 93-103), cloned vascular smooth muscle cells (Suttles et al., 1995, Exp. Cell Res. 218: 331-338), endothelial cell-derived chemoattractant exposed T cells (Katz et al., 1994, J. Leukoc. Biol. 55: 567-573), cultured human aortic endothelial cells (Farber et al., 1992, Am. J. Physiol. 262: H1088-1085), and foam cell cultures (Libby et al., 1996, Curr Opin Lipidol 7: 330-335). Potentially effective Therapeutics, for example but not by way of limitation, reduce foam cell formation in cell culture models, or reduce atherosclerotic plaque formation in hypercholesterolemic mouse models of atherosclerosis in comparison to controls.

Accordingly, once an atherosclerosis-associated disease or disorder has been shown to be amenable to treatment by modulation of activity or formation, that disease or disorder can be treated or prevented by administration of a Therapeutic that modulates activity.

Cytokine and Cell Proliferation/Differentiation Activity

A GENX protein of the present invention may exhibit cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may induce production of other cytokines in certain cell populations. Many protein factors discovered

to date, including all known cytokines, have exhibited activity in one or more factor dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of a protein of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+ (preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e and CMK.

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The activity of a protein of the invention may, among other means, be measured by the following methods: Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by Coligan et al., Greene Publishing Associates and Wiley-Interscience (Chapter 3 and Chapter 7); Takai et al., J Immunol 137:3494-3500, 1986; Bertagnoili et al., J Immunol 145:1706-1712, 1990; Bertagnolli et al., Cell Immunol 133:327-341, 1991; Bertagnolli, et al., J Immunol 149:3778-3783, 1992; Bowman et al., J Immunol 152:1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described by Kruisbeek and Shevach, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan *et al.*, eds. Vol 1, pp. 3.12.1-14, John Wiley and Sons, Toronto 1994; and by Schreiber, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan eds. Vol 1 pp. 6.8.1-8, John Wiley and Sons, Toronto 1994.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described by Bottomly et al., In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto 1991; deVries et al., J Exp Med 173:1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc Natl Acad Sci U.S.A. 80:2931-2938, 1983; Nordan, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.6.1-5, John Wiley and Sons, Toronto 1991; Smith et al., Proc Natl Acad Sci U.S.A. 83:1857-1861, 1986; Measurement of human Interleukin 11-Bennett, et al. In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto 1991; Ciarletta, et al., In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and

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cytokine production) include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds., Greene Publishing Associates and Wiley-Interscience (Chapter 3Chapter 6, Chapter 7); Weinberger et al., Proc Natl Acad Sci USA 77:6091-6095, 1980; Weinberger et al., Eur J Immun 11:405-411, 1981; Takai et al., J Immunol 137:3494-3500, 1986; Takai et al., J Immunol 140:508-512, 1988.

Immune Stimulating or Suppressing Activity

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A GENX protein of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A protein may be useful in the treatment of various immune deficiencies and disorders (including severe combined immunodeficiency (SCID)), e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by vital (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases causes by vital, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpesviruses, mycobacteria, Leishmania species., malaria species. and various fungal infections such as candidiasis. Of course, in this regard, a protein of the present invention may also be useful where a boost to the immune system generally may be desirable, i.e., in the treatment of cancer.

Autoimmune disorders which may be treated using a protein of the present invention include, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitus, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein of the present invention may also to be useful in the treatment of allergic reactions and conditions, such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein of the present invention.

Using the proteins of the invention it may also be possible to immune responses, in a number of ways. Down regulation may be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response.

The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or energy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon re-exposure to specific antigen in the absence of the tolerizing agent.

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Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions (such as, for example, B7), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a molecule which inhibits or blocks interaction of a B7 lymphocyte antigen with its natural ligand(s) on immune cells (such as a soluble, monomeric form of a peptide having B7-2 activity alone or in conjunction with a monomeric form of a peptide having an activity of another B lymphocyte antigen (e.g., B7-1, B7-3) or blocking antibody), prior to transplantation can lead to the binding of the molecule to the natural ligand(s) on the immune cells without transmitting the corresponding costimulatory signal. Blocking B lymphocyte antigen function in this matter prevents cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, the lack of costimulation may also be sufficient to energize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of B lymphocyte antigens.

The efficacy of particular blocking reagents in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the

immunosuppressive effects of CTLA4Ig fusion proteins *in vivo* as described in Lenschow *et al.*, Science 257:789-792 (1992) and Turka *et al.*, Proc Natl Acad Sci USA, 89:11102-11105 (1992). In addition, murine models of GVHD (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of blocking B lymphocyte antigen function *in vivo* on the development of that disease.

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Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and auto-antibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block costimulation of T cells by disrupting receptor: ligand interactions of B lymphocyte antigens can be used to inhibit T cell activation and prevent production of auto-antibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythematosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 840-856).

Upregulation of an antigen function (preferably a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may be in the form of enhancing an existing immune response or eliciting an initial immune response. For example, enhancing an immune response through stimulating B lymphocyte antigen function may be useful in cases of viral infection. In addition, systemic vital diseases such as influenza, the common cold, and encephalitis might be alleviated by the administration of stimulatory forms of B lymphocyte antigens systemically.

Alternatively, anti-viral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells *in vitro* with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the *in vitro* activated T cells into the

patient. Another method of enhancing anti-vital immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells *in vivo*.

In another application, up regulation or enhancement of antigen function (preferably B lymphocyte antigen function) may be useful in the induction of tumor immunity. Tumor cells (e.g., sarcoma, melanoma, lymphoma, leukemia, neuroblastoma, carcinoma) transfected with a nucleic acid encoding at least one peptide of the present invention can be administered to a subject to overcome tumor-specific tolerance in the subject. If desired, the tumor cell can be transfected to express a combination of peptides. For example, tumor cells obtained from a patient can be transfected ex vivo with an expression vector directing the expression of a peptide having B7-2-like activity alone, or in conjunction with a peptide having B7-1-like activity and/or B7-3-like activity. The transfected tumor cells are returned to the patient to result in expression of the peptides on the surface of the transfected cell. Alternatively, gene therapy techniques can be used to target a tumor cell for transfection in vivo.

The presence of the peptide of the present invention having the activity of a B lymphocyte antigen(s) on the surface of the tumor cell provides the necessary costimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient amounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I α chain protein and β_2 microglobulin protein or an MHC class II a chain protein and an MHC class II β chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor

specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

The activity of a protein of the invention may, among other means, be measured by the following methods: Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Greene Publishing Associates and Wiley-Interscience (Chapter 3, Chapter 7); Herrmann et al., Proc Natl Acad Sci USA 78:2488-2492, 1981; Herrmann et al., J Immunol 128:1968-1974, 1982; Handa et al., J Immunol 135:1564-1572, 1985; Takai et al., J Immunol 137:3494-3500, 1986; Takai et al., J Immunol 140:508-512, 1988; Herrmann et al., Proc Natl Acad Sci USA 78:2488-2492, 1981; Herrmann et al., J Immunol 135:1564-1572, 1985; Takai et al., J Immunol 137:3494-3500, 1986; Bowman et al., J Virology 61:1992-1998; Takai et al., J Immunol 140:508-512, 1988; Bertagnolli et al., Cell Immunol 133:327-341, 1991; Brown et al., J Immunol 153:3079-3092, 1994.

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Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, *J Immunol* 144:3028-3033, 1990; and Mond and Brunswick In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan *et al.*, (eds.) Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto 1994.

Mixed lymphocyte reaction (MLR) assays (which will identify, among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan *et al.*, eds. Greene Publishing Associates and Wiley-Interscience (Chapter 3, Chapter 7); Takai *et al.*, *J Immunol* 137:3494-3500, 1986; Takai *et al.*, *J Immunol* 140:508-512, 1988; Bertagnolli *et al.*, *J Immunol* 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J Immunol 134:536-544, 1995; Inaba et al., J Exp Med 173:549-559, 1991; Macatonia et al., J Immunol 154:5071-5079, 1995; Porgador et al., J Exp Med 182:255-260, 1995; Nair et al., J Virol 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., J Exp Med 169:1255-1264, 1989; Bhardwaj et al., J Clin Investig 94:797-807, 1994; and Inaba et al., J Exp Med 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Res 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, J Immunol 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., Internat J Oncol 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cell Immunol 155: 111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc Nat Acad Sci USA 88:7548-7551, 1991.

Hematopoiesis Regulating Activity

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A GENX protein of the present invention may be useful in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell deficiencies. Even marginal biological activity in support of colony forming cells or of factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines, thereby indicating utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either in-vivo or ex-vivo (i.e., in conjunction with bone marrow

transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson *et al. Cellular Biology* 15:141-151, 1995; Keller *et al.*, *Mol. Cell. Biol.* 13:473-486, 1993; McClanahan *et al.*, *Blood* 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, In: Culture of Hematopoietic Cells. Freshney, et al. (eds.) Vol pp. 265-268, Wiley-Liss, Inc., New York, N.Y 1994; Hirayama et al., Proc Natl Acad Sci USA 89:5907-5911, 1992; McNiece and Briddeli, In: Culture of Hematopoietic Cells. Freshney, et al. (eds.) Vol pp. 23-39, Wiley-Liss, Inc., New York, N.Y. 1994; Neben et al., Exp Hematol 22:353-359, 1994; Ploemacher, In: Culture of Hematopoietic Cells. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, N.Y. 1994; Spoonceret al., In: Culture of Hematopoietic Cells. Freshhey, et al., (eds.) Vol pp. 163-179, Wiley-Liss, Inc., New York, N.Y. 1994; Sutherland, In: Culture of Hematopoietic Cells. Freshney, et al., (eds.) Vol pp. 139-162, Wiley-Liss, Inc., New York, N.Y. 1994.

Tissue Growth Activity

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A GENX protein of the present invention also may have utility in compositions used for bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as for wound healing and tissue repair and replacement, and in the treatment of burns, incisions and ulcers.

A protein of the present invention, which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Such a preparation employing a protein of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation

induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A protein of this invention may also be used in the treatment of periodontal disease, and in other tooth repair processes. Such agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of progenitors of bone-forming cells. A protein of the invention may also be useful in the treatment of osteoporosis or osteoarthritis, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes.

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Another category of tissue regeneration activity that may be attributable to the protein of the present invention is tendon/ligament formation. A protein of the present invention, which induces tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions of the present invention may provide an environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendonitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a career as is well known in the art.

The protein of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, *i.e.* for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a protein

may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nervous injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a protein of the invention.

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Proteins of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

It is expected that a protein of the present invention may also exhibit activity for generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine, kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring to allow normal tissue to regenerate. A protein of the invention may also exhibit angiogenic activity.

A protein of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A protein of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for tissue generation activity include, without limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without limitation, those described in: Winter, EPIDERMAL WOUND HEALING, pp. 71-112 (Maibach and Rovee, eds.), Year Book Medical

Publishers, Inc., Chicago, as modified by Eaglstein and Menz, J. Invest. Dermatol 71:382-84 (1978).

Activin/Inhibin Activity

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A GENX protein of the present invention may also exhibit activin- or inhibin-related activities. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a protein of the present invention, alone or in heterodimers with a member of the inhibin a family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the protein of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin-b group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, U.S. Pat. No. 4,798,885. A protein of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as cows, sheep and pigs.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc Natl Acad Sci USA 83:3091-3095, 1986.

Chemotactic/Chemokinetic Activity

A protein of the present invention may have chemotactic or chemokinetic activity (e.g., act as a chemokine) for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. Chemotactic and chemokinetic proteins can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic proteins provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example,

attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

The activity of a protein of the invention may, among other means, be measured by following methods:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: Current Protocols in Immunology, Coligan et al., eds. (Chapter 6.12, Measurement of Alpha and Beta Chemokines 6.12.1-6.12.28); Taub et al. J Clin Invest 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller et al., Eur J Immunol 25: 1744-1748; Gruberet al. J Immunol 152:5860-5867, 1994; Johnston et al., J Immunol 153: 1762-1768, 1994.

Hemostatic and Thrombolytic Activity

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A protein of the invention may also exhibit hemostatic or thrombolytic activity. As a result, such a protein is expected to be useful in treatment of various coagulation disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A protein of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as, for example, infarction of cardiac and central nervous system vessels (e.g., stroke).

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res.

45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

Receptor/Ligand Activity

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A protein of the present invention may also demonstrate activity as receptors, receptor ligands or inhibitors or agonists of receptor/ligand interactions. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell—cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses). Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: CURRENT PROTOCOLS IN IMMUNOLOGY, Ed by Coligan, et al., Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc Natl Acad Sci USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J Immunol Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

Anti-Inflammatory Activity

Proteins of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell—cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Proteins exhibiting such activities can be used to treat inflammatory conditions including chronic or acute

conditions), including without limitation inflammation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Proteins of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material.

Tumor Inhibition Activity

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In addition to the activities described above for immunological treatment or prevention of tumors, a protein of the invention may exhibit other anti-tumor activities. A protein may inhibit tumor growth directly or indirectly (such as, for example, via ADCC). A protein may exhibit its tumor inhibitory activity by acting on tumor tissue or tumor precursor tissue, by inhibiting formation of tissues necessary to support tumor growth (such as, for example, by inhibiting angiogenesis), by causing production of other factors, agents or cell types which inhibit tumor growth, or by suppressing, eliminating or inhibiting factors, agents or cell types which promote tumor growth.

Other Activities

A protein of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape); effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, cofactors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders) and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting

deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

Neural disorders in general include Parkinson's disease, Alzheimer's disease, Huntington's disease, multiple sclerosis, amyotrophic lateral sclerosis (ALS), peripheral neuropathy, tumors of the nervous system, exposure to neurotoxins, acute brain injury, peripheral nerve trauma or injury, and other neuropathies, epilepsy, and/or tremors.

EQUIVALENTS

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From the foregoing detailed description of the specific embodiments of the invention, it should be apparent that particular novel compositions and methods involving nucleic acids, polypeptides, antibodies, detection and treatment have been described. Although these particular embodiments have been disclosed herein in detail, this has been done by way of example for purposes of illustration only, and is not intended to be limiting with respect to the scope of the appended claims that follow. In particular, it is contemplated by the inventors that various substitutions, alterations, and modifications may be made as a matter of routine for a person of ordinary skill in the art to the invention without departing from the spirit and scope of the invention as defined by the claims. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying figures. Such modifications are intended to fall within the scope of the appended claims.



Table 1

*000	ODE# Intomal Identification	Destain eimitarity			Language Transfer of the Country of the
# LVO	Number				Calls of dissues in which cene is Expressed
-	13076366 (1, 2)	Novel Protein sim. GBank gil4691395 emb CAB41562.1 - (AL049727) putative large secreted protein [Streptomyces coelicolor]			264636
2	80248091 (3, 4)	Novel Protein sim. GBank gil2829506[splP71559]SUCC_MYCTU - SUCCINYL-COA SYNTHETASE BETA CHAIN (SCS-BETA)	Contains protein domain (PF00549) - UNCLASSIFIED CoA-ligases		264907, 264600, 264602, 264762, 264769, 264689, 264638, 264567
ဇ	80415924 (5, 6)				264910, 264604, 264634, 264905, 264636, 264691, 264907, 264692, 264629
4	82018837 (7, 8)			UNCLESSIFIED	264908, 264909, 264760, 264628, 264635
5	79970035 (9, 10)				22279002, 264563
ယ	79842462 (11, 12)		Contains protein domain (PF00127) - UNCLASSIFIED Conser binding proteins		264908
			γ		
- ω	56924278 (15, 16)	Novel Protein sim. GBank Ularabidopsis thaliana] Novel Protein sim. GBank Ggi585562 sp[006458 NINB_KLEPN - NITRITE GGI585562 sp[006458 NINB_KLEPN - NITRITE GGI585658 NDD/PN) ARGE SIIRI INIT		reductase	264905, 264906, 264907, 264909, 264909, 264906, 264906, 264908, 264909, 264908, 264909, 264908, 264909, 264906, 264512, 265009, 264909, 264595, 264596, 264761, 265009, 264094, 264760, 264693, 264689, 35695917, 264690, 264692, 264693, 33657109, 264628, 264629, 35696423, 264634, 264636, 264636, 264639, 18108385, 264563, 264564, 264566, 264907
6	79394457 (17, 18)			Π	265007, 265019, 263972
, <u>e</u>	79556459 (19, 20)			UNCLASSIFIED	264906
=	20414027 (21, 22)				264605
12	94141210 (23, 24)	Novel Protein sim. GBank gij3878145 emb CAA99871 - (Z75543) similar to potassium channel protein [Caenorhabditis elegans]			264259, 265007, 83373044
13	20750551 (25, 26)				264556, 264557, 264564
4	95105114 (27, 28)	Novel Protein sim. GBank gilz832781 emb CA412645 - Contains pr (AJ225805) inward potassium channel alpha subunit [Egeria Ank repeat densa]	Contains protein domain (PF00023) - potassium_channel Ank repeat		35696286, 35696052, 264510, 35695917, 264691, 264628, 35696423, 264555, 264558, 264559, 83373044
15	20458307 (29, 30)	Novel Protein sim. GBank gij1710791[sp]Q10234[RT05_SCHPO - PROBABLE MITOCHONDRIAL 40S RIBOSOMAL PROTEIN S5	Contains protein domain (PF00333) - irbosomalprot Ribosomal protein S5		264604
16	20760356 (31, 32)				264555

17	20292744 (33, 34)	Novel Protein sim. GBank	Contains protein domain (PF00449) -		264600
		gij1174884 sp P44391 URE1_HAEIN - UREASE ALPHA SUBUNIT (UREA AMIDOHYDROLASE)	Urease		
18	80246804 (35, 36)	Novel Protein sim. GBank gi[2281102 (AC002333) - SF16 isolog [Arabidopsis thaliana]			29331827, 264555, 264557, 264638, 264558
19	80076624 (37, 38)			UNCLASSIFIED	22278996, 264907, 264910, 264600, 264693
20	20724558 (39, 40)	Novel Protein sim. GBank		transport	264602
		gi 2506112 sp P43672 UUP_ECOLI - ABC TRANSPORTER ATP-BINDING PROTEIN UUP			
21	80417554 (41, 42)	Novel Protein sim. GBank		UNCLASSIFIED	22278995, 264906, 265008, 265010, 265011,
		gij1730203jspjP50442jGATM_RAT - GLYCINE			264602, 264605, 264766, 264688, 21906764,
_		AMIDINOTRANSFERASE PRECURSOR (L-			264691, 18108376, 264636, 18108387,
		ARGININE:GLYCINE AMIDINOTRANSFERASE)			264486
1	11705858 (43 44)				264685
16		Norm Protein eim CBank eilse77220lambiC&B070771	Contains and demain (DE00441) debudrosesses	dobudooooo	264488 264007 264000 264600 264602
3	80419175 (45, 46)	Novel Protein sim. Gbank gij 1877/329jemoj CABU7077 - 72027777 (adE25 (Macabadarium tuberculosis)	Contains protein domain (Proud41)	genyarogenase	204460, 204907, 204909, 204800, 204002, 264601 264605 264682 264766 32833986
		לבסדים ליין מסרבים ליין כספספופיותייי ותספו כתוספו			264636, 264486
	20291697 (47, 48)				264600
25	80253774 (49, 50)				264593
1	80255394 (51, 52)			UNCLASSIFIED	22278996, 56182435, 265018, 264566
ı	80235795 (53, 54)	Novel Protein sim. GBank gil4808369lemb[CAB42783.1] -	Contains protein domain (PF00253) - inbosomatprot	ribosomalprot	18108370, 35696423, 264635, 264555
		(AL049841) putative 30S ribosomal protein S14	Ribosomal protein S14p/S29e		
ą	70483584 /55 5E)	loughes menonal		INCLASSIEIED	25463B
3	(22, 22)			20000	מנטרטט טטברטט טטברטט רפטרטס
₹	82448765 (57, 58)	Novel Protein sim. GBank gij3122290 sp 008333 K6PF_STRCO - 6- PHOSPHOFRUCTOKINASE (PHOSPHOFRUCTOKINASE) (PHOSPHOHEXOKINASE) (ATP-PFK)	Contains protein domain (PT 00365) - Kinase Phosphofructokinase	Kinase	.204001, 204 <i>1</i> 02, 204 <i>1</i> 00, 204 <i>1</i> 09, 204050
ణ	79189333 (59, 60)			UNCLASSIFIED	264908, 265019, 264687, 21906764, 21906766
31	19848158 (61, 62)			UNCLASSIFIED	264534
32	82449495 (63, 64)	Novel Protein sim. GBank gij3560504 (AF027770) - unknown [Mycobacterium smegmatis]		UNCLASSIFIED	264905, 264605, 264762, 264766, 264687, 264689
ន	79582628 (65, 66)	Novel Protein sim. GBank gi[2129003]pirj G64507 - hypothetical protein MJ1665 - Methanococcus jannaschii		UNCLASSIFIED	264687
<u>청</u>	87467657 (67, 68)			UNCLASSIFIED	60432289, 264600, 264602, 264760, 18108357, 264769, 265020, 264691
32	95005170 (69, 70)	Novel Protein sim. GBank gi[5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	264600, 264687, 264558, 264639
36	19642042 (71, 72)	Novel Protein sim. GBank gij3287739 sp p73538 BIOB_SYNY3 - BIOTIN SYNTHASE (BIOTIN SYNTHETASE)		synthase	264566
37	20369215 (73, 74)	Novel Protein sim. GBank gil2313134 gb AAD07126.11 - (AE000527) delta-1-pyrroline-5-carboxylate dehydrogenase [Helicobacter pylori 26695]		dehydrogenase	264603

264605	264905, 264906, 264907, 66712502, 264908, 264909, 264511, 265009, 264910, 55812038, 264758, 265011, 264762, 264682, 264763, 264764, 264631, 264633, 264631, 264631, 264631, 264631, 264536, 18108385, 264482	264592	264591, 35695917	264602	264605	264769, 264636	264769, 264510, 264508	264566	264689	18108385, 264635, 264828	264603	264508, 264603, 264769, 264689, 264636, 264558, 264486	264593, 18108387	264634	264762	264630, 264909, 264766	29331824, 264102, 265018, 18108376	264604	264557	
reductase	complement	UNCLASSIFIED		UNCLASSIFIED	synthase	synthase	UNCLASSIFIED	ngf	UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED			UNCLASSIFIED	
	Contains protein domain (PF00207) - complement Alpha-2-macroglobulin family					Contains protein domain (PF00958) - synthase GMP synthase C terminal domain													Contains protein domain (PF00076) - UNCLASSIFIED	RBD, or RNP domain)
Novel Protein sim. GBank gij3805970jembjCAA062311 - (AJ004933) periplasmic nitrate reductase, large subunit IRhodooseudomonas sp. l	Novel Protein sim. GBank gi[1929449 (L63543) - endodermin [Xenopus laevis]		Novel Protein sim. GBank gij854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]		Novel Protein sim. GBank gij3820584 (AF086791) - carbamoyiphosphate synthelase large subunit [Zymomonas mobilis]	Novel Protein sim. GBank gi[2494784]sp Q50729 GUAA_MYCTU · GMP SYNTHASE (GLUTAMINE-HYDROLYZING) (GLUTAMINE AMIDOTRANSFERASE) (GMP SYNTHETASE)	Novel Protein sim. GBank gil 1881738 (U89688) - myosin-l binding protein Acan125 (Acanthamoeba castellanii)						Novel Protein sim. GBank gij3411177 (AF076240) - MocC [Rhizoblum leguminosarum bv. viciae]	Novel Protein sim. GBank gij3914992[sp]Q26264 SM41_HEMPU - 41 KD SPICULE MATRIX PROTEIN PRECURSOR (HSM41) (HPSMC)	Novel Protein sim. GBank gil3980411 (AC004561) - putative protine-rich protein [Arabidopsis thaliana]		Novel Profein sim. GBank gil 1633572 (U52064) - Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma- associated herpes-like virus]	Novel Protein sim. GBank gil4321580[gb[AAD15785] - (AF050114) alqinate Iyase [Pseudomonas sp. W7]		
20466334 (75, 76)	94300715 (77, 78)	20635625 (79, 80)	80023287 (81, 82)	20724566 (83, 84)	20467069 (85, 86)	13085297 (87, 88)	39384711 (89, 90)	95003398 (91, 92)	11698624 (93, 94)	79407218 (95.96)	21659844 (97, 98)	80503996 (99, 100)	80255569 (101, 102)	79208528 (103, 104)	36996970 (105, 106)	79570897 (107, 108)	80202703 (109, 110)	8758408 (111, 112)	11223386 (113, 114)	
38	39	40	2	42	43	4.	45	46	2 2	- 4	T	S	51	52	53	5	25	29	25	

8	91227306 (113, 110)	Novel Protein sim. GBank gi 5616074 gb AAD45616.1 AF06194 - (AF061943) protate-	Contains protein domain (PFUUU59) - kinase Eukaryotic protein kinase domain	Kinase	56182575, 264259, 60432049, 35696052. 66712502, 264909, 265008, 265010, 265011.
		derived STE20-like kinase PSK (Homo sapiens)			264681, 29148784, 35695917, 60170615.
					264691, 264692, 264693, 18108374,
					35696423, 56182323, 60432113
29	80077371 (117, 118)	Novel Protein sim. GBank	Contains protein domain (PF00953) - Irransferase	transferase	264600, 264689, 264638
		gi 1172920 sp P45830 RFE_MYCLE - PUTATIVE	Glycosyl transferase		
		UNDECAPRENYL-PHOSPHATE ALPHA-N-			
		ACETYLGLUCOSAMINYLTRANSFERASE			
00 00	12958341 (119, 120)				264689
<u>6</u>	80426806 (121, 122)	Novel Protein sim. GBank gi[1710216 (U79260) - unknown		glycoprotein	264766
		[Homo sapiens]			
62	13504966 (123, 124)			;	264630
63	16474553 (125, 126)			UNCLASSIFIED	265019
2	20724578 (127, 128)	Novel Protein sim. GBank gil420945 pir A47041 -		UNCLASSIFIED	264602
		inseri			
		Alcaligenes eutrophus			
92	79326308 (129, 130)	Novel Protein sim. GBank	Contains protein domain (PF00224) - kinase	kinase	264563
		gij3122312 sp O06134 KPYK_MYCTU - PYRUVATE	Pyruvate kinase		
		KINASE (PK)			
99	46854384 (131, 132)	Novel Protein sim. GBank gij3928723 emb CAA22219 -		transport	22278996, 264558
		(AL034355) putative ABC transporter (Streptomyces			
		coelicotor			
29	78952543 (133, 134)	Novel Protein sim. GBank		dehydrogenase	265021
	•	Gil231985IspIP30234IDHA MYCTU - ALANINE		1	
		DEHYDROGENASE (40 KD ANTIGEN)			
68	79817382 (135, 136)				264909
69	79841764 (137, 138)			UNCLASSIFIED	264908
20	79871329 (139, 140)				264906, 264908
71	65897456 (141, 142)			UNCLASSIFIED	264602, 265021
12	87734977 (143 144)	Novel Protein sim GBank oil4415926lphtAAD201571		INC. ASSIFIED	26448B 264905 264906 264907 264908
<u>. </u>		(ACO05282) unknown protein [Arabidonsis thaliana]			264511 265008 264910 264758 87168474
					264682 264766 264686 264689 35695917
					265021 E0170615 264691 33657023
					20021, 00170310, E01031, 0001 0E01
					22279000
73	80025241 (145, 146)			UNCLASSIFIED	60424179, 264508, 264908, 265007, 264603,
					264687, 264689, 264692, 18108387
74	20377410 (147, 148)			UNCLASSIFIED	264605
75	11819032 (149, 150)	Novel Protein sim. GBank gil2853098 emb CAA16914 -		UNCLASSIFIED	264689
		(AL021767) vacuolar protein sorting [Schizosaccharomyces pombe]			
76	95105303 (151, 152)	Novel Protein sim. GBank gi[4468811 emb CAB38212 -		UNCLASSIFIED	83373044, 264906, 264557
		(AL035601) putative protein [Arabidopsis thaliana]			
۲.	10144718 (153, 154)	Novel Protein sim. GBank gi 854065 emb CAA58337 - (X83413) UB8 [Human herpesvirus 6]		UNCLASSIFIED	264563
28	8758258 (155, 156)			UNCLASSIFIED	264604

62	94140190 (157, 158)	Novel Protein sim. GBank qil5689453idbilBAA83010.11 -	Contains protein domain (PF00169) -		3550628E 22278008 20331822 20331824
		(AB028981) KIAA1058 protein [Homo sapiens]	PH domain		29331825, 29331827, 264905, 264906
					264907, 66712502, 264908, 264909, 265008.
					265009, 264910, 60170831, 55812038,
					33109954, 265017, 265018, 264288, 264768,
					56181562, 21906765, 21906769, 29148784,
					265020, 264690, 264691, 264692, 264693.
					60431528, 35696423, 264631, 264632,
					264634, 264636, 264639, 83373044, 264564,
3					264566, 264567
⊋	82314840 (158, 160)			UNCLASSIFIED	264769, 264601, 265006, 264910, 264604,
					264605, 264634, 264635, 264905, 264762.
					264637, 264592, 264628, 264907, 264691,
إ					264908, 264567, 264909, 264766
<u>.</u>	20467247 (161, 162)	Novel Protein sim. GBank		reductase	264605
		gij1723442lsplQ10258lYD2A_SCHPO - HYPOTHETICAL			
		69.0 KD PROTEIN C56F8.10 IN CHROMOSOME I			
82	16331388 (163, 164)	Novel Protein sim. GBank gi 2895866 (AF045770) -		dehydrogenase	264567
		methylmalonate semi-aldehyde dehydrogenase (Oryza			
		satival			
83	94741180 (165, 166)	Novel Protein sim. GBank gi[3402673 (AC004697) -		UNCLASSIFIED	264488, 264508, 264509, 264905, 264908.
		unknown protein [Arabidopsis thaliana]			264909 264511 264591 264593 264594
					264505 264506 264759 264603 264360
					204030, £04030, £04700, £04700, £04700,
					204061, 10100331, 264/62, 264682, 264/64,
					264684, 264766, 264686, 264632, 264637,
إ	1000				264557, 264638, 264639, 18108385, 264566
ž	80355375 (167, 168)			transport	264508, 264906, 264907, 264908, 264909,
		gij11/3364 sp P45380 SAT1_RAT - SULFATE ANION			264910, 284760, 264763, 264764, 264766,
	·	TRANSPORTER 1 (CANALICULAR SULFATE			264768, 264769, 35695855, 264636, 264637
		TRANSPORTER) (SULFATE/CARBONATE ANTIPORTER)			
82	80499600 (169, 170)	Novel Protein sim. GBank gil2120998[pir] S70682 -		transferase	264605, 264762, 264687, 264769, 18108374,
					264636, 264486
8	39559043 (171, 172)	Novel Protein sim. GBank gij3256023jemb/CAA17228.1f -			264910
		(AL021897) hypothetical protein Rv1112 (Mycobacterium			
		[tuberculosis]			
87	13856808 (173, 174)			UNCLASSIFIED	264093

88	85344718 (175, 176)	Novel Protein sim. GBank gil559703 dbj BAA07552 - (D38549) ha 1025 is new [Homo sapiens]			52644507, 52646365, 18108398, 65274572, 56182575, 56994075, 35696286, 22278997,
					22278998, 22278999, 264092, 264093, 264094, 264095, 264096, 264096, 264096, 264097, 26331822, 29331824, 56182181, 66714117, 29331825, 29331824, 56182181, 66714117, 29331826, 29331826, 29331826, 29331826, 29351827, 29331826, 264906, 264906, 264907, 264006, 264908, 29331830, 265009, 264906, 264906, 264907, 264906, 264907, 264906, 264907, 264907, 264906, 264907, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 265011, 87168559, 264601, 265017, 265018, 264066, 265019, 26448, 264369, 264286, 21906766, 21906767, 265020, 265021, 265022, 60170615, 26504150, 33657039, 35695709, 264628, 263972, 18108374, 55810764, 3659423, 5611576, 652747891, 35695855, 60170394, 83373044, 18108388, 18108388, 56526486, 87168518, 60432113, 25279000, 22279000, 264486, 264486
68	80077389 (177, 178)	Novel Protein sim. GBank gij 1710383 sp P46352 RIPX_BACSU - PROBABLE INTEGRASE/RECOMBINASE RIPX			264600
8	82115999 (179, 180)			UNCLASSIFIED	264760
16	78906950 (181, 182)	Novel Protein sim. GBank gi 2499891 sp P76403 YEGQ_ECOLI • PUTATIVE PROTEASE IN BAER-OGRK INTERGENIC REGION	a.		265006
85	79554871 (183, 184)	Novel Protein sim. GBank gil3367754[emb[CAA20079] - (AL031155) hypothetical protein SC3A7.16c [Streptomyces conjication]		UNCLASSIFIED	264691
83	80496778 (185, 186)	Novel Protein sim. GBank gij2895095 (AF011337) - putative E1-E2 ATPase (Mus musculus)		ATPase_associated	ATPase_associated 264907, 264908, 264910, 265009, 264605, 264769
3	79646649 (187, 188)	A BACSU - GLYCINE BETAINE	Contains protein domain (PF00571) - transport	ransport	264906
95	11090238 (189, 190)				264594

8 	i_	Novel Protein sim. GBank gil4589560jdbj BAA76802.1 j - (AB023175) KIAA0958 protein [Homo sapiens]			22278995, 22278999, 264259, 29331822, 29331826, 35696052, 29146499, 244509, 264906, 264907, 264908, 264909, 264909, 265009, 265008, 264910, 265009, 264593, 265010, 265017, 264604, 265019, 18108351, 264288, 264766, 264769, 21906765, 21906767, 21906769, 265020, 264629, 33657182, 35695763, 264628, 18108379, 264631, 264636, 18108381, 264559, 18108382, 83373044, 22279002
97	79605200 (193, 194)	Novel Protein sim. GBank giļ4583559jemb CAB40388.1 - [(AJ005255) OxyR [Erwinia chrysanthemi]		UNCLASSIFIED	264508
88	79427000 (195, 196)	Novel Protein sim. GBank gi 1001693 dbj BAA10430 - (D64002) hypothetical protein [Synechocystis sp.]		UNCLASSIFIED	264909
66	20466524 (197, 198)	Novel Protein sim. GBank gij1169479jspjP43925jEFG_HAEIN - ELONGATION FACTOR G (EF-G)		UNCLASSIFIED	264605
5	79640113 (199, 200)			UNCLASSIFIED	264693
5	80203298 (201, 202)			UNCLASSIFIED	265020, 264102, 263972
102	20467259 (203, 204)	Novel Protein sim. GBank gi 2894166 emb CAA11773.1 - (AJ223998) PCZA361.18 [Amycolatopsis orientalis]		synthase	264605
103	20466368 (205, 206)	Novel Protein sim. GBank gil1731040 sp P54509 YQHH_BACSU - HYPOTHETICAL HELICASE IN SINI-GCVT INTERGENIC REGION	Contains protein domain (PF00271) - helicase Helicases conserved C-terminal domain		264605
호	80247572 (207, 208)	Novel Protein sim. GBank gil854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264591, 264595, 264602
105	79605206 (209, 210)	Novel Protein sim. GBank gi 1685117 (U70770) - furrowed [Drosophila melanogaster]	Contains protein domain (PF00084) - complement Sushi domain (SCR repeat)		264508
99	28382058 (211, 212)	Novel Protein sim. GBank gi 1705505 sp P54729 BS4_MOUSE - BS4 PROTEIN	Contains protein domain (PF00527) - UBA domain	UNCLASSIFIED	264511, 265009
107	80057791 (213, 214)	Novel Protein sim. GBank gil4887229[gb AAD32244.1 AF15075 - (AF150755) microtubule-actin crosslinking factor [Mus muscutus]		associated	ATPase_associated 29331824, 264591, 21906754, 265019
108	80237936 (215, 216)	Novel Protein sim. GBank gi 2635771 emb CAB15264 - (299120) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis]	Contains protein domain (PF00005) - Iransport ABC transporter		18108374, 35695917, 22278996, 264113, 264600, 264602, 264603, 285017, 264910, 264906, 264636, 264766
109	95194148 (217, 218)	Novel Protein sim. GBank gi[2330791]emb[CAB11265] - (298601) carboxypeptidase s precursor [Schizosaccharomyces pombe]		UNCLASSIFIED	264758, 264603, 264630, 264636, 264637
110	79582823 (219, 220)				264687
111	39565458 (221, 222)				264564
112	79856038 (223, 224)			١	264908
	17959439 (225, 226)			UNCLASSIFIED	265007
=	80502101 (227, 228)			1	264769

133	10887507 (763 264)				000100
2 5	1000/002 (203, 204)				264636
3	94030003 (203, 200)	Nover Protein sim. Cbank gij1877340jembjcAb070b8j - (292771) accA3 [Mycobacterium tuberculosis]	Contains protein domain (PF00289) - carboxylase Carbamoyl-phosphate synthase (CPSase)	carboxylase	264905, 264689
<u>동</u>	79834660 (267, 268)	Novel Protein sim. GBank gil4585838 emb CAB40932.1 - (AL049630) putative NADH dehydrogenase [Streptomyces coelicolor]		dehydrogenase	264905, 264605, 265021
135	19885057 (269, 270)	Novel Protein sim, GBank gil1460074 emb CAB01049 - (277250) hypothetical protein Rv2566 [Mycobacterium [tuberculosis]			264634
136	79846083 (271, 272)	Novel Protein sim. GBank gi 2125896 emb CAA73511 - (Y13070) folylpolyglutamate synthase [Streptomyces coellcolor]		synthase	264508
137	79619770 (273, 274)				264683, 264685, 264686, 264691, 264692, 264693
138	79635971 (275, 276)	Novel Protein sim. GBank gil5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	18108374, 18108385, 33657109, 33657182, 265010, 22278998, 265006, 265007, 265008, 265009, 264693
139	86688076 (277, 278)	Novel Protein sim. GBank gij5689912 emb CAB52075.1 - (AL109732) putative mutase [Streptomyces coelicolor A3(2)]	Contains protein domain (PF01817) - dehydrogenase Chorismate mutase	dehydrogenase	22278996, 265007, 264910, 60433356, 265010, 264602, 264605, 264766, 264688, 264769, 264693, 32833986, 18108374, 18108387
140	79825759 (279, 280)			UNCLASSIFIED	264908
141	20700094 (281, 282)				264600
142	80028104 (283, 284)	Novel Protein sim. GBank gij3581916 emb CAA20855 - (AL031545) mutS family DNA mismatch repair protein [Schizosaccharomyces pombe]			264602, 265017
143	11072274 (285, 286)			UNCLASSIFIED	264600
<u>4</u>	95009102 (287, 288)	Novel Protein sim. GBank gij3334127[sp P97303]BAC2_MOUSE - TRANSCRIPTION REGULATOR PROTEIN BACH2 (BTB AND CNC HOMOLOG 2)			263978, 264600, 264910, 264632, 264508, 264563, 264564, 264591, 264556, 264908, 264629, 264639
145	80027058 (289, 290)	Novel Protein sim. GBank gil3757569[emb[CAA21315] - (AL031863) 1-evidence=predicted by content; 1-method=genefinder;084; 1-method_score=66.31; 1-evidence_end [Drosophila melanogaster]		UNCLASSIFIED	22278996, 264602
146	13085662 (291, 292)	Novel Protein sim. GBank gil140807[sp]P24536]YI21_BURCE - INSERTION ELEMENT IS402 HYPOTHETICAL 24 KD PROTEIN	Contains protein domain (PF01675) - Transposase		264687
147	94320366 (293, 294)	Novel Protein sim. GBank gil2827608 emb CAA16663 - (AL021646) uvrD2 [Mycobacterium tuberculosis]		helicase	264905, 264906, 264909, 264510, 265009, 60433356, 264600, 284601, 264604, 264605, 264687, 264769, 18108365, 65274791, 18108387
148	80248804 (295, 296)	Novel Protein sim. GBank gi 2916947 emb CAA17585 - (AL021999) hypothetical protein Rv0986 [Mycobacterium [tuberculosis]		transport	265009, 265010, 264600, 264602, 264603, 264604, 264605, 264693, 33657109, 264636

149	IR0249373 (297 298)	Novel Protein cim CBank	Table of the state	7	
		gi1723073 sp Q11040 7081_MYCTU - HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN CYS0.01	ABC transporter		2030 iu, 2040uu, 2040u i, 2040u3, 2040u4, 27486265, 264636
150	20294748 (299, 300)	Novel Protein sim. GBank gi 3724125 emb CAA11905 - (AJ224340) maltosephosphorylase [Lactobacillus			264600
		sanfrancisco]			
151	20726398 (301, 302)	Novel Protein sim. GBank qi1729312IspIP07651IDEOB ECOLL-	Contains protein domain (PF01676) - UNCLASSIFIED Metalloenzome connectamily	UNCLASSIFIED	264602
		PHOSPHOPENTOMUTASE (PHOSPHODEOXXRIBOMI TASE)			
152	95002877 (303, 304)	Novel Protein sim. GBank		pentidase	264602
		gi[2497952[sp P55667 74TM_RHISN - HYPOTHETICAL HYDROLASE/PEPTIDASE Y4TM			
153	80256665 (305, 306)	Novel Protein sim. GBank		UNCLASSIFIED	264593
		gij3123021 splQ90508 VIT1_FUNHE - VITELLOGENIN I			
	· · · · · · · · · · · · · · · · · · ·	PRECURSOR (VIG.1) (CONTAINS: LIPOVITELLIN 1 (LV1); PHOSVITIN (PV); LIPOVITE(1) N 2 (1 V2))			
154	82305966 (307, 308)				264910, 264762, 264691, 264634
155	20429859 (309, 310)	Novel Protein sim. GBank gil419697/pir/JJN0443 -	Contains protein domain (PF00140) - Inapolymerase	rnapolymerase	264605
			Sigma-70 factor		
		Streptomyces aureofaciens			
156	39564742 (311, 312)	Novel Protein sim. GBank gij628710 pir S41739 -		UNCLASSIFIED	264565
157	10358887 (313, 314)	Novel Protein sim. GBank gij3695013 (AF052586) - CtrA	Contains protein domain (PF00142) - hydrolase	hydrolase	264691
		[Pseudomonas aeruginosa]	4Fe-4S iron sulfur cluster binding		
450	1010 000 0000		proteins, NifH/frxC family		
90	1970 1930 (313, 310)	Nover Protein sim. GBank gij 1073072 prij C55543 - cmaU protein - Pseudomonas syringae pv. syringae		UNCLASSIFIED	264905
159	78890376 (317, 318)			UNCLASSIFIED	265008
9	11075119 (319, 320)		Contains protein domain (PF00400) -		264605
;	1000 1000 1000		WD domain, G-beta repeat		
191	80055007 (321, 322)	Novel Protein sim. GBank gi[1173023]sp P46789 RL30_STRCO - 50S RIBOSOMAL PROTEIN L30	Contains protein domain (PF00327) - Inbosomalprot Ribosomal protein L30p/L7e	ribosomalprot	22278996, 264600, 264603, 35695917, 32833986, 35696423, 264636
162	80016371 (323, 324)	Novel Protein sim. GBank gil5304869 emb CAB46028.1 -	Contains protein domain (PF00097) - interleukin	interleukin	264112, 264532, 22279002
		(ALUS 1665) dJ963KZS.Z (novel prolein) [Homo sapiens]	Zinc tinger, C3HC4 type (KING finaer)		
163	11692306 (325, 326)			j	264639
164	80077902 (327, 328)			UNCLASSIFIED	264905, 264907, 264600
165	10856067 (329, 330)				264691
166	88095003 (331, 332)	Novel Protein sim. GBank gi 2661691 emb CAA15795 - (AL009204) putative protease [Streptomyces coelicolor]		UNCLASSIFIED	264605, 264486
167	16395460 (333, 334)	Novel Protein sim. GBank giļ4416478 gb AAD20378 - (AF125999) transposase {Mycobacterium avium}		UNCLASSIFIED	265010
	80079362 (335, 336)	Novel Protein sim. GBank gil76177 Ipir IQQECFT - hypothetical 38.8K protein (ffs! 5' region) - Escherichia coli			264600
169	80239581 (337, 338)				264556, 264557, 264558, 264559

170	79612364 (339, 340)				307730
171	95293073 (341, 342)	Novel Protein sim. GBank gl 140888 sp P27847 YIGK_ECOLI - HYPOTHETICAL 15.4 KD PROTEIN IN RECQ-PLDB INTERGENIC REGION {F138}	Contains protein domain (PF01810) - LysE type translocator		264595, 264604
172	37797007 (343, 344)	Novel Protein sim. GBank gil4210905 gb AAD12048.1 - (AF045609) AgIG [Sinorhizoblum meliloti]	Contains protein domain (PF00528) - transport Binding-protein-dependent transport systems inner membrane component		264769
173	57529660 (345, 346)	Novel Protein sim. GBank gi 132854 sp P02387 RL2_ECOL Contains protein domain (PF00181) - nbosomatprot - 50S RIBOSOMAL PROTEIN L2	Contains protein domain (PF00181) - Ribosomal Proteins L2	ribosomaiprot	264769
174	95293078 (347, 348)	Novel Protein sim. GBank gi 1881350 db BAA19377 - (AB001488) PROBABLE TRANSPORT PROTEIN, SIMILAR TO ANTIBIOTIC TRANSPORT-ASSOCIATED PROTEIN ACTII IN STREPTOMYCES COELICOLOR. [Bacillus subtilis]		transport	264510, 264593, 264602, 264603, 264605, 264762, 264693
175	79756270 (349, 350)	Novel Protein sim. GBank gij2072722jemb CAB08326 - (295121) manA [Mycobacterium tuberculosis]		isomerase	264565
176	80066896 (351, 352)	Novel Protein sim. GBank gil1055198 (U40187) - similar to PIR:A41724 chicken LD (limb deformity) gene product and to formin; also P-rich region similar to collagen [Caenorhabditis elegans]		UNCLASSIFIED	264907, 264910, 264681, 264558
14	86684852 (353, 354)	Novel Protein sim. GBank gi 2326738 emb CAB10952 - (298268) hypothetical protein Rv1695 [Mycobacterium tuberculosis]	Contains protein domain (PF01513) - UNCLASSIFIED Domain of unknown function		264768, 60424179, 264687, 264688, 264769, 29331826, 60432289, 18108376, 264689, 18108387, 32833986, 22278996, 265020, 264600, 264601, 264602, 264603, 264604, 264605, 264638, 264638, 264562, 264564, 2646637, 264638, 264486, 60433356, 264766
178	79559526 (355, 356)	Novel Protein sim. GBank gil 1906596 (U81788) - kinesin-73 [Drosophila melanogaster]		struct	264693, 33657109, 264635
179	20263112 (357, 358)			UNCLASSIFIED	264563
180	80488958 (359, 360)	Novel Protein sim. GBank gi 1169367 sp P45256 DNAB_HAEIN - REPLICATIVE DNA HELICASE		helicase	264769
181	79585369 (361, 362)	Novel Protein sim. GBank gij3170615 (AF059485) - DOC4 [Mus musculus]		UNCLASSIFIED	21906767, 264635, 264639, 18108384
182	80577899 (363, 364)			UNCLASSIFIED	264259, 35696052, 56182435, 264511, 265018, 33657109, 264555, 264566
183	11614017 (365, 366)	Novel Protein sim. GBank gi 1076627 pir S54172 - inorganic pyrophosphatase (EC 3.6.1.1) - common tobacco		UNCLASSIFIED	264690
184	10174167 (367, 368)	Novel Protein sim. GBank gil4371280[gb]AAD18138] - (AC006260) hypothetical protein [Arabidopsis thaliana]		UNCLASSIFIED	264510

21660822 (369, 370)	Novel Protein sim. GBank gil3006178 emb CAA18398.1 - (AL022304) putative mma transport regulator Schizosaccharomyces pombel		UNCLASSIFIED	264604
80070329 (371, 372)	Novel Protein sim. GBank gij2829802 sp P94408 YCLF_BACSU - HYPOTHETICAL 53.3 KD PROTEIN IN SFP-GERKA INTERGENIC REGION		transport	264595
80186611 (373, 374)			UNCLASSIFIED	264369
20464942 (375, 376)	Novel Protein sim. GBank gij3150260 emb CAA19179 - [AL023634] cyclin [Schizosaccharomyces pombel		kinase	264605
82338215 (377, 378)	Novel Protein sim. GBank gij2145853 pir S72938 - hflX protein - Mycobacterium leprae		UNCLASSIFIED	35696052, 264602, 264605, 264762, 264689, 35695917, 18108370, 18108372, 264638, 264565
80086821 (379, 380)	Novel Protein sim. GBank gil 1881244 dbj BAA19271 - (AB001488) SIMILAR TO PYRUVATE OXIDASE AND ACETOLACTATE SYNTHASE. [Bacillus subtilis]	Contains protein domain (PF00205) - synthase Thiamine pyrophosphate enzymes	synthase	264563
88095012 (381, 382)	Novel Protein sim. GBank gi[120226 sp P28725 FKBP_STRCH - FK506-BINDING PROTEIN (PEPTIDYL-PROLYL CIS-TRANS ISOMERASE) (PPIASE) (ROTAMASE)	Contains protein domain (PF00254) - isomerase FKBP-type peptidyl-protyl cis-trans isomerases	somerase	264508, 264604, 264605, 264769, 264555
16333379 (383, 384)				264567
79910127 (385, 386)				264908, 264693
20464949 (387, 388)				264605
13518389 (389, 390)	Novel Protein sim. GBank gil4980892 gb AAD35474.1 AE00171 - (AE001718) ABC transporter, ATP-binding protein [Thermotoga maritima]		transport	264636
95005569 (391, 392)	Novel Protein sim. GBank gi[1705461]splp53656 BIOA_ERWHE - ADENOSYLMETHIONINE-8-AMINO-7-OXONONANOATE AMINOTRANSFERASE (7.8-DIAMINO-PELARGONIC ACID AMINOTRANSFERASE) (DAPA AMINOTRANSFERASE)	Contains protein domain (PF00202) - gaba Aminotransferases dass-III pyridoxal phosphate	jaba	264600, 264689, 264638
80248665 (393, 394)	Novel Protein sim. GBank gi[3122305 sp Q27778 K6PF_SCHMA - 6- PHOSPHOFRUCTOKINASE (PHOSPHOFRUCTOKINASE) (PHOSPHOHEXOKINASE)	Contains protein domain (PF00365) - kinase Phosphofructokinase	tinase	264602, 264662, 264692, 18108374
79163635 (395, 396)				264636
78890715 (397, 398)	Novel Protein sim. GBank gi 1781203 emb CAB06110 - (283859) gnd [Mycobacterium tuberculosis]	Contains protein domain (PF00393) - 6-phosphogluconate dehydrogenases		265008
79413849 (399, 400)	Novel Protein sim. GBank gi 2642222 (AF030885) - telomere-associated recQ-like helicase [Ustilago maydis]		UNCLASSIFIED	264595, 264596
86945924 (401, 402)	Novel Protein sim. GBank gij2894379jembjCAA74911.1j. (Y14573) ring finger protein [Hordeum vulgare]		UNCLASSIFIED	29331826, 265007, 264512, 33657402, 264596, 265017, 18108351, 264682, 264683, 264767, 264629, 55810764, 264634, 264635, 56182323, 60432113, 22279000

202	70598045 (402 404)	News Order of the Control			
3	(tot 'cot)	Nover Frotein stri. GBank gil231772 sp P30598 CHS1_USTMA - CHITIN SYNTHASE 1 (CHITIN-UDP ACETYL-GLUCOSAMINYL TRANSFERASE 1)	Contains protein domain (PF01644) - synthase Chilin synthase	synthase	264600
203	79843927 (405, 406)	Novel Protein sim. GBank gi 1504042 dbj BAA13220 - (D86984) similar to yeast adenylate cyclase (S56776) Homo sapiens			22278995, 29331822, 29331825, 29331827, 264906, 21906754, 264683, 21906766, 34606473, 364664
204	79855186 (407, 408)			INCI ASSIFIED	2564909
205	10090583 (409, 410)	Novel Protein sim. GBank gij2633808 emb CAB13310] - (Z99111) similar to hypothetical proteins [Bacillus subtilis]			264909
206	8758473 (411, 412)			UNCLASSIFIED	264604
207	20754522 (413, 414)	Novet Protein sim. GBank gi 2134381 pir S60678 - polybrome 1 protein - chicken			284556
208	20289261 (415, 416)			1	264605
209	80071069 (417, 418)	Novel Protein sim. GBank			264605, 264689
		gi 2501040 sp 005814 SYP_MYCTU - PROLYL-TRNA SYNTHETASE (PROLINETRNA LIGASE) (PRORS)			
210	80168800 (419, 420)				264905, 264907, 264909, 264766, 264687, 264691, 264629, 18108374, 264638
211	80034539 (421, 422)				263978
212	82442474 (423 424)	Novel Protein sim CRank		G1131334 13141	00100 001000 001000 001000
	(12)	gi 5031809 ref NP_005536.1 pISLR - immunoglobulin superfamily containing leucine-rich repeat		UNCLASSIFIED	.264508, 264905, 264907, 264908, , , 264608, , , , 264600, 264762, 264634, 264635, 264639, 264635, 264636, , , , , , , , , , , , , , , , , , ,
213	80249562 (425, 426)	Novel Protein sim. GBank	Contains protein domain (PF00330) - isomerase		22278996, 264508, 264600, 264602, 264603,
		gij3122359jspj033123 LEU2_MYCLE - 3- ISOPROPYLMALATE DEHYDRATASE LARGE SUBUNIT (ISOPROPYLMALATE ISOMERASE) (ALPHA-IPM	Aconitase family (aconitate hydratase)		264605, 33657023, 264565, 264486
	1007 1007 1000	ISOMERASE) (IPMI)			
<u>4</u>	(900/9361 (42/, 426)	Novel Protein sim. GBank gi[116238[sp]P19421[CH60_COXBU - 60 KD CHAPERONIN (PROTEIN CPN60) (GROEL PROTEIN) (HEAT SHOCK PROTEIN B)	Contains protein domain (PF00118) - eph TCP-1/cpn60 chaperonin famity		264600, 264693
215	14973283 (429, 430)			0	264629
216	80177716 (431, 432)	Novel Protein sim. GBank gij3417297 (AC002310) - Unknown gene product [Homo sapiens]	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type		264448
217	79603634 (433, 434)	Novel Protein sim. GBank gil2506924 sp P49754 VP41_HUMAN - VACUOLAR ASSEMBLY PROTEIN VPS41 HOMOLOG (S53)			264508
218	80258475 (435, 436)	Novel Protein sim. GBank gij1173288jspjP38106jRSEA_ECOLI - SIGMA-E FACTOR NEGATIVE REGULATORY PROTEIN		rnapolymerase	264594
219	20438797 (437, 438)	Novel Protein sim. GBank gi 1781097 emb CAB06231 - (283864) gilB [Mycobacterium tuberculosis]		synthase	264604
220	13499572 (439, 440)	Novel Protein sim. GBank gil2984703 (AF052427) - unknown [Trypanosoma cruzi]			264689
221	11287498 (441, 442)	Novel Protein sim. GBank giļ4587313(dbj BAA76709.1 - (AB025248) alpha-1,2-mannosidase [Bacillus sp. M-90]		UNCLASSIFIED	264555

222	79862802 (443, 444)	Novel Protein sim. GBank gi1877268jemb CAB07049j - (292770) hypothetical protein Rv0143c [Mycobacterium tubercutosis]		UNCLASSIFIED	284605, 264769, 35696423
223	83053869 (445, 446)			UNCLASSIFIED	264906, 264907, 264603
777	79557920 (447, 448)				264684, 264693
225	79559541 (449, 450)	Novèl Protein sim. GBank gi[2274851 dbj BAA21515 - (D64159) 3-7 gene product [Homo sapiens]		UNCLASSIFIED	264692
226	79172397 (451, 452)	Novel Protein sim. GBank gi 868245 (U29488) - C56C10.7 gene product Caenomabditis elegans		UNCLASSIFIED	22278998, 264112, 33657023, 263981
227	81777196 (453, 454)			UNCLASSIFIED	35695917, 264636, 264907
228	79872285 (455, 456)				264768, 264907, 264908, 264692, 264593, 264639
229	79838266 (457, 458)				264906, 264910
230	11013209 (459, 460)			UNCLASSIFIED	264631
231	20622207 (461, 462)	Novel Protein sim. GBank gi 1835114 emb CAA71733 - (Y10744) homoserine O-acetyltransferase [Leptospira meyen]			264906, 264600, 264603, 264692
232	80055035 (463, 464)			UNCLASSIFIED	264600, 264603, 264605, 264687, 264769
233	80063054 (465, 466)	Novel Protein sim. GBank gij2642340 (AF032970) - imidazolone propionate hydrolase (Pseudomonas putida)	Contains protein domain (PF00449) - hydrotase Urease	hydrolase	264604
234	7523998 (467, 468)	Novel Protein sim. GBank gij3510505 (AF030881) - pol polyprotein (Firau nibrioes)		UNCLASSIFIED	264369
35	80203671 (469 470)			OBIBION ION	064406
318	7004004 (474 470)			UNCLASSIFIED	204 100
8	78940001 (471, 472)	Novel Protein sim. GBank gil2104609 emb CAB08805 - (295398) PckA [Mycobacterium leprae]		carboxylase	264905
2	11755273 (473, 474)				264681
238	79461401 (475, 476)			UNCLASSIFIED	264639
239	82435190 (477, 478)		Contains protein domain (PF00037) -		264906, 265010, 264603, 264762, 264682,
		gi[2495617[sp]Q57252[YOJJ_HAEIN - HYPOTHETICAL PROTEIN HI1163	4Fe-4S ferredoxins and related ironsulfur cluster binding domains.		264636, 264638, 264486
240	21635575 (479, 480)	Novel Protein sim. GBank gij3183458 sp P75796 YLIA_ECOLI - HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN YLIA		transport	264259, 264769
241	80377307 (481, 482)	Novel Protein sim. GBank gij3875920jemb[CAB04111] - (Z81503) predicted using Genefinder; similar to collagen; CDNA EST EMBL:D65450 comes from this gene; cDNA EST EMBL:D68888 comes from this gene [Caenorhabditis]		UNCLASSIFIED	264908, 264909, 264764, 264639
242	82148454 (483, 484)			UNCLASSIFIED	264489, 264907, 264908, 264511, 264760, 264764, 264692, 264635, 264637
243	79633207 (485, 486)			UNCLASSIFIED	264906
244	80248682 (487, 488)	Novel Protein sim. GBank gi 2624302 emb CAA15575 - (AL008967) ald [Mycobacterium tuberculosis]		dehydrogenase	264600, 264602, 264605, 264769, 264689
245	79863543 (489, 490)	Novel Protein sim. GBank gi 2920625 (AF044499) - vgrE protein [Escherichia coli]		UNCLASSIFIED	264907, 264758
246	79162929 (491, 492)	Novel Protein sim. GBank gi[5420387 emb CAB46679.1] - (AJ243459) proteophosphoglycan [Leishmania major]	Contains protein domain (PF01106) - NifU-like domain		264637, 18108381, 18108387, 264565

247	79873185 (493, 494)	Novel Protein sim. GBank gij 1839006jembjCAB06648j -		kinase	264909, 264691, 35696423, 18108387
		(Z85982) argB (Mycobacterium tuberculosis)			
248	80488983 (495, 496) 	Novel Protein sim. GBank gi 1168574 sp P42464 ATPB_CORGL - ATP SYNTHASE BETA CHAIN		synthase	35696286, 264907, 264511, 264602, 264768. 264688, 265021, 35695855, 18108385
249	79764645 (497, 498)			UNCLASSIFIED	264907, 264910, 265011, 264762, 264636
250	79619980 (499, 500)				21906768, 264692
251	84359489 (501, 502)			UNCLASSIFIED	52645156, 29331822, 29331824, 52644045, 265018, 21906765, 21906768, 265020, 27466261, 27486265, 35695763, 18108376, 264556, 264556
252	79737756 (503, 504)	Novel Protein sim. GBank gij3327166 dbj BAA31651 - (AB014576) KIAA0676 protein [Homo sapiens]			264685, 264687, 264632
253	20443124 (505, 506)	Novel Protein sim. GBank gij3036880[emb CAA18513] - (AL022374) putative ATP-dependent DNA helicase (Streptomyces coelicolor)		helicase	264604
254	80027421 (507, 508)	Novel Protein sim. GBank gi[3915488 sp 034961 YJMB_BACSU - HYPOTHETICAL SYMPORTER IN COTT-RAPA INTERGENIC REGION		UNCLASSIFIED	264508, 264906, 264602, 264687, 265021, 264486
255	11398315 (509, 510)	9 8		UNCLASSIFIED	264593
256	80028158 (511, 512)	Novel Protein sim. GBank gil465787 sp P34422 YL31_CAEEL - HYPOTHETICAL 86.0 Prolyl oligopeptidase family KD PROTEIN F44B9.1 IN CHROMOSOME III	Contains protein domain (PF00326) - peptidase Prolyl oligopeptidase family	peptidase	264602, 264692
257	20289282 (513, 514)	Novel Protein sim. GBank gi[1172039]sp[P42315]SCOA_BACSU - PROBABLE SUCCINYL-COA:3-KETOACID-COENZYME A TRANSFERASE SUBUNIT A (SUCCINYL COA:3-OXOACID COA-TRANSFERASE) (OXCT A)	Contains protein domain (PF01144) - transferase Coenzyme A transferase	transferase	264605
258	20459464 (515, 516)	Novel Protein sim. GBank gij3127836 emb CAA18902 - (AL023496) hypothetical protein (Streptomyces coelicolor)		UNCLASSIFIED	264604
529	79910152 (517, 518)				264681, 264686, 264692
097	20379437 (519, 520)				264692, 264556
192	20285883 (521, 522)	Novel Protein sim. GBank gi 123761 sp P24221 HUTH_STRGR - HISTIDINE AMMONIA-LYASE (HISTIDASE)	Contains protein domain (PF00221) - UNCLASSIFIED Phenylalanine and histidine ammonia lyases		264600
262	80189317 (523, 524)			UNCLASSIFIED	265017, 264369
263	88095045 (525, 528)	Novel Protein sim. GBank gil3924708 emb CAA84646 - (Z35597) Weak similarity with sea squirt nidogen precursor protein (blastp score 71); cDNA EST EMBL:102069 comes from this gene; cDNA EST EMBL:076135 comes from this gene; cDNA EST EMBL:073147 comes from this gene; cDNA EST EMBL:073147 comes from this gene;		UNCLASSIFIED	264488, 264905, 264906, 264907, 264908, 264909, 264512, 264910, 264758, 264596, 264604, 26519, 264605, 264760, 18108351, 264763, 264764, 284288, 264768, 284768, 264768, 264631, 264634, 264635, 264635, 264635, 264638, 264638,
264	87370826 (527, 528)	Novel Protein sim. GBank gi[3043734 dbj BAA25531 - (AB011177) KIAA0605 protein [Homo sapiens]	Contains protein domain (PF00047) - protease Immunoglobulin domain	protease	264259, 264908, 21906754, 265018, 265019, 265020

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265	95355646 (529, 530)	Novel Protein sim. GBank gil4589624 dbj BAA/6834.1 ABN232071 KIAA0990 nrolein [Homo saniens]	Kinase	254468, 35696269, 29331624, 50162181, 35696052, 264508, 264905, 264906, 264907,
				66712502, 264908, 264909, 264511, 264512,
				264910, 264592, 264595, 264758, 264596,
				55811386, 264600, 265017, 264603, 264604,
				264605, 264760, 18108351, 264762, 264681,
				264764, 264288, 264766, 264768, 264769,
				21906765, 21906767, 21906769, 265020,
				264691, 33657023, 33657109, 33657182,
				264628, 35696423, 35695855, 264630,
				264631, 264632, 264634, 264635, 264636,
				264555, 264638, 83373044, 56526486,
				87168518, 264564, 264566, 264486
266	79588075 (531, 532)			264600
267	11362222 (533, 534)		UNCLASSIFIED	264828
268	79909566 (535, 536)		UNCLASSIFIED	264687, 264769, 264689
269	80025810 (537, 538)		UNCLASSIFIED	264602
270	84361144 (539 540)	Novel Protein sim. GBank		264693
<u> </u>		gil4507367refiNP_003182.1pTARS - threonyl-tRNA		
		synthetase		
271	79552301 (541, 542)		UNCLASSIFIED	264909, 264693
272	9574778 (543, 544)	Novel Protein sim. GBank	synthase	264908
		gil4980738 gb AAD35331.1 AE00170 - (AE001707) glucose-		
		1-phosphate adenylyltransferase [Thermotoga maritima]		
273	12840694 (545, 546)	Novel Protein sim. GBank	UNCLASSIFIED	264688
		gi[1168224 sp P44569 5NTO_HAEIN - PROBABLE 5'- INICI FOTIDASE PRECURSOR		
274	39524246 (547, 548)			264564
275	82787041 (549, 550)	Novel Protein sim. GBank gij3253159 (AF005355) -	UNCLASSIFIED	264907, 264908, 264909, 264766, 264768,
		Irranslation initiation factor eIF2C (Oryctolagus cuniculus)		264691, 264632, 264636
276	86671073 (551, 552)	Novel Protein sim. GBank		265008, 60432229
		gij134920ispjP21997iSSGP_VOLCA - SULFATED SLIREACE GLYCOPROTFIN 185 (SSG 185)		
277	80079735 (553, 554)	Novel Protein sim. GBank	ribosomalprot	264600, 18108387
		gi[129021 sp P20964 OBG_BACSU - SPO08-ASSOCIATED GTP-BINDING PROTEIN		
278	12966947 (555, 556)		UNCLASSIFIED	264689
278	95282719 (557, 558)	Novel Protein sim. GBank gil79839 pir S03812 - uvrB	nuclease	264508, 264604, 21906764, 264638, 264557,
		protein - Micrococcus luteus		264404
280	5603617 (559, 560)			264259
281	80249599 (561, 562)	Novel Protein sim. GBank		18108392, 264634, 264555, 264556, 264557,
		gi3123160jspjQ18964 YLN2_CAEEL - HYPOTHETICAL		264558
		45.2 KD I KP-ASP REPEALS CONTAINING PROTEIN D2013.2 IN CHROMOSOME II		
282	18598682 (563, 564)		UNCLASSIFIED	265019
283	20614211 (565, 566)		UNCLASSIFIED	264555

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284	91212160 (567, 568)		Contains protein domain (PF00300) - UNCLASSIFIED	UNCLASSIFIED	35696052, 29331828, 264508, 264905,
		[xylan esterase; AxeA [Thermotoga neapolitana]	Phosphoglycerate mutase family		264600, 264602, 264605, 264682, 264764,
					56181562, 21906764, 18108376, 264636,
					264559, 18108387
285	8757940 (569, 570)			UNCLASSIFIED	264603
286	80503235 (571, 572)	Novel Protein sim. GBank gi 2072674 emb CAB08305 -	Contains protein domain (PF00270) - ATPase_associated 35696052, 264769, 264638	ATPase_associated	35696052, 264769, 264638
		(Z95120) rhiE [Mycobacterium tuberculosis]	DEAD/DEAH box helicase		
287	12745521 (573, 574)			UNCLASSIFIED	264689
288	20756502 (575, 578)	Novel Protein sim. GBank gil765323 bbs 157676 - (S74439)		collagen	264557
		silk fibroin heavy chain (C-terminal) [Bombyx			
		mori=silkworms, Peptide Partial, 633 aaj [Bombyx mori]			
289	80043804 (577, 578)	Novel Protein sim. GBank gil 1870009 emblCAB068601 -	Contains protein domain (PF00440) - ribosomalprot	ribosomalprot	264593, 264600
		(Z92539) hypothetical protein Rv1019 (Mycobacterium	Bacterial regulatory proteins, tetR		
		[uberculosis]	family		
280	80430175 (579, 580)			UNCLASSIFIED	264768
291	20747431 (581, 582)	Novel Protein sim. GBank		UNCLASSIFIED	264601
		ail2506664ispiP40120IYDCG ECOLI - 59.4 PROTEIN IN			
292	80052555 (583, 584)	Novel Protein sim. GBank gi 625182 (L39015) -		UNCLASSIFIED	264605
_		mitochondrial glutamyl-tRNA synthelase [Saccharomyces			
		cerevisiae]			
293	80062519 (585, 586)	Novel Protein sim. GBank		helicase	264909, 264605, 264687, 264689, 264692
_		gil1718065 spiP53528 UVRD_MYCLE - PUTATIVE DNA			
		HELICASE II HOMOLOG			
584	79830303 (587, 588)	Novel Protein sim. GBank	Contains protein domain (PF00008) - oncogene	oncogene	35696052, 264906, 265011, 264628,
		gi 117422 sp P10040 CRB_DROME - CRUMBS PROTEIN PRECURSOR (95F)	EGF-like domain		55811576
282	79444180 (589, 590)	Novel Protein sim. GBank gil1181619jdbj BAA11565 -			52644507, 29331822, 264592, 265020,
		(D82364) a variant of TSC-22 [Gallus gallus]			264639
967	79607076 (591, 592)	Novel Protein sim. GBank gil3649789 dbj BAA33403 -		synthase	264508
		(AB012226) SecA [Vibrio alginolyticus]			
297	79631297 (593, 594)	Novel Protein sim. GBank gil5689967 lemblCAB52004.11 -		UNCLASSIFIED	264905, 264687, 264638
		(AL109663) putative membrane protein (Streptomyces			
- 1		coelicolor A3(2)]			
298	80418898 (595, 596)			UNCLASSIFIED	264905, 264691, 264639, 264766

66 7	95283298 (597, 598)	Novel Protein sim. GBank gil220637[dbj BA401477] - (D10627) zinc finger protein [Mus musculus]	Contains protein domain (PF00096) Zinc finger, C2H2 type		264486, 263994, 56994075, 22278997, 22278998, 22278999, 20221099, 29331824, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264905, 264906, 264907, 264908, 2644045, 264908, 264906, 264907, 264908, 2644045, 264908, 264908, 264908, 264768, 26408, 265019, 264768, 264687, 56181562, 264769, 264687, 56181562, 264769, 264692, 33657109, 27486261, 18108370, 264628, 264629, 55811576, 35695855, 264631, 264634, 264635, 264639, 26279000,
8	20711340 (599, 600)			INCI ASSIFIED	22279002, 264565, 264566, 264567 264602
301	13511332 (601, 602)	Novel Protein sim. GBank gil145922 (M20981) - iron dictirate transport protein precursor [Escherichia coli]		transport	264687
302	9875260 (603, 604)	Novel Protein sim. GBank gi[1174661[sp]P44594]TGT_HAEIN - QUEUINE TRNA- RIBOSYLTRANSFERASE (TRNA-GUANINE TRANSGLYCOSYLASE) (GUANINE INSERTION ENZYME)			264908
303	79574895 (605, 606)				264689
304	20711344 (607, 608)	Novel Protein sim. GBank gil67985 pirj HJNVAV - helicase (EC 3.6.1) - Autographa californica nuclear polyhedrosis virus		helicase	284602
305	80412520 (609, 610)	Novel Protein sim. GBank gij728867 sp P40602 APG_ARATH - ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR			264763
306	8515876 (611, 612)	Novel Protein sim. GBank gil1657554 jgbJAAB18082.1 - (U73857) hypothetical protein [Escherichia coli]		UNCLASSIFIED	263978
307	80222901 (613, 614)			UNCLASSIFIED	265010, 21906768, 265020, 18108374, 263977
308	80064305 (615, 616)	Novel Protein sim. GBank gil1710612 sp Q10793 RNH2_MYCTU - PROBABLE RIBONUCLEASE HII (RNASE HII)	Contains protein domain (PF01351) - nuclease Ribonuclease HII	nuclease	264910, 264600, 264605, 264687, 264689, 264638, 18108387
309	80504136 (617, 618)	Novel Protein sim. GBank gil5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]			264769
310	80053616 (619, 620)				264603
311	11090659 (621, 622)	Novel Protein sim. GBank gi 1144522 (U34957) - phosphoribosylaminoimidazolesuccinocarboxamide synthase [Mycobacterium tuberculosis]	<u>.</u>	synthase	264602
312	80054347 (623, 624)			UNCLASSIFIED	264566
313	80046168 (625, 626)				264603, 264567

	07045440 (007 000)	" " " " " " " " " " " " " " " " " " "			
	020, 021, 020)	(Nover Protein Sint. Godin gijaco (acost (Acost (2) - Ikaros (Danjo renio)	GATA zinc finger	dna_ena_end	284239, 60432289, 29331828, 264903, 284906 284908 284909 285008 284910
					60432229 33657402 60433438 33109954
					265011, 265017, 264603, 265018, 264288,
_					264766, 264692, 35695763, 264628, 264629,
					264639, 60170394, 22279002, 264568
315	82356091 (629, 630)	Novel Protein sim. GBank gil 1652620 dbj BAA17540 -			264508, 264600, 264762, 264687, 264768,
		(D90907) pyridine nucleotide transhydrogenase beta			52644229, 264769, 264689, 264635, 264636,
9,0	1000 1001 1001	subunit jaynechocystis sp.j			264638, 264486
	789 107 (631, 632)			UNCLASSIFIED	264693
317	(20466944 (633, 634)			UNCLASSIFIED	264605
		<pre> gi 118244 sp P24176 DAPE_ECOLI - SUCCINYL-</pre>		,	
		DIAMINOPIMELATE DESUCCINYLASE (SDAP)			
318	94141836 (635, 636)	Novel Protein sim. GBank	Contains protein domain (PF00526) - Itransport	transport	264908, 264909, 264910, 264593, 264594,
		gij4680229jgbjAAD27583.1JAF11827 - (AF118274) DNb-5	Dictyostelium (slime mold) repeats	•	264760, 264288, 264768, 264769, 21906769,
		[Homo sapiens]			264691, 264693, 264628, 65274791, 264635,
					264636, 264638, 83373044, 22279002.
					264566
319	17289360 (637, 638)	Novel Protein sim. GBank gil1149693 emb CAA60220 -		transport	265018
		-			
320	13527675 (639, 640)			svnthase	264687
_		gi[2811033]sp O05314 GLGC MYCTU - GLUCOSE-1-			
		PHOSPHATE ADENYLYI TRANSFERASE (ADP.			
		GLUCOSE SYNTHASE) (ADP-GLUCOSE			
		PYROPHOSPHORY (ASE)			
321	94134387 (641, 642)	Novel Protein sim. GBank qi 1680716 (U68234) - all-trans-		cvto450	264509, 264906, 264907, 264908, 265009.
		retingic acid 4-hydroxylase (Danio rerio)			DEARDE DRATEA DEARDE DEARDA DEARDE
					264638 264639 83373044 264567
33	GEARONS (EAS)	Novel Protein cim CBack ail1160355 (1133059) 1 INC 90		OF ACCIENCE	ECO114ED DEAED1 ED121FD EED107E1
356	(cro, crt)	Nover Floreit Sint. Spain, 9 1,003333 (033030) - 0140-03 [Caenorhabditis elegans]		ONCCASSIFIED	556 1 50, 2040 1, 8043 526 556 10/64
323	94653725 (645, 646)			UNCLASSIFIED	264488, 265009, 264593, 264628, 264635
324	(79174383 (647, 648)				264687
325	79862691 (649, 650)			UNCLASSIFIED	264693
328	28774974 (651, 652)			UNCLASSIFIED	264288, 18108385
327	79776267 (653, 654)	Novel Protein sim. GBank gil451544 (U04267) - proline-rich cell wall protein [Gossypium barbadense]			264488, 264905, 264509, 264910
328	80253202 (655, 656)			UNCLASSIFIED	264592
329	10173821 (657, 658)			UNCLASSIFIED	264510
330	86597767 (659, 660)	Novel Protein sim. GBank gil4191358 (AF087825) - claudin-		UNCLASSIFIED	264259, 264908
		7 [Mus musculus]			
331	79754888 (661, 662)	Novel Protein sim. GBank ail807411pirlIS20912 - regulatory		transcriptfactor	264910 264687 264689 264636 264567
		protein whiB - Streptomyces coeffcolor			
332	80071440 (663, 664)			reductase	35696423, 264636, 264638, 264565
		gij114049jspjP19480jAHPF_SALTY - ALKYL			
		HYDROPEROXIDE REDUCTASE SUBUNIT F (ALKYL			
500	1999 300: 1110000	HYDROPEROXIDE REDUCTASE F52A PROTEIN)			
333	13009555 (665, 666)				264687

264905, 284600, 264604, 264486	264907, 264603, 264693, 18108374, 264636, 18108387	265009, 264766, 264686	264602	264569, 18108397, 22278998, 29331822, 20281099, 29331824, 56162181, 66714117, 29331825, 35696052, 29331828, 264508, 264509, 264906, 264906, 264907, 264908, 264909, 265008, 264900, 265008, 264901, 265019, 265019, 264768, 55812038, 6527444, 265011, 264785, 264681, 264762, 264762, 18108351, 264682, 2644229, 264699, 55811957, 35895917, 264693, 264639, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264639, 18108376, 18108380, 264638, 264638, 264639, 18108380, 264638, 264638, 264639, 18108389, 183373044, 18108385, 87168518, 60432113	264593	264905, 265019, 264769, 18108374	264605	264603, 264604	264605	264600, 264605, 264768, 18108370, 18108374, 35695655	264593	264905
dehydrogenase	esterase	UNCLASSIFIED		nucl_recpt	eph	ubiquilin	isomerase			UNCLASSIFIED	histone	dehydrogenase
Contains protein domain (PF00208) - Glutamate/Leucine/Phenylalanine/Valine dehydrogenase				Contains protein domain (PF00249) - Inucl_recpt Myb-like DNA-binding domain			Contains protein domain (PF00290) - isomerase Tryptophan synthase alpha chain					Contains protein domain (PF00208) - dehydrogenase Glutamate/Leucine/Phenylalanine/Va line dehydrogenase
n n	Novel Protein sim. GBank gi[2193938]emb[CAB09602] - (296800) gipQ2 [Mycobacterium tuberculosis]			Novel Protein sim. GBank gil5454074 ref NP_006303.1 pSMRT - silencing mediator for Myb-like DNA-binding domain retinoid and thyroid hormone receptors	Novel Protein sim. GBank gil4001713 dbj BAA35087.1 - (AB015879) DnaK [Porphyromonas gingivalis]	Novel Protein sim. GBank gitz842699[sp]Q92353]UBPC_SCHPO - PUTATIVE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE C6G9.08 (UBIQUITIN THIOLESTERASE) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING	Novel Protein sim. GBank gij2686580 (AE001166) - conserved hypothetical protein Borrelia burgdorferij	Novel Protein sim. GBank gi 1684738 emb CAA70601 - (Y09452) Yed hypothetical protein [Pseudomonas syringae]	5	Novel Protein sim. GBank gil2117275[emb[CAB09104] - (295618) hypothetical protein Rv0807 [Mycobacterlum Inberculosis]	Novel Protein sim. GBank gi]3023317[sp]048935[aPHA_MYCRA - ACETYLPOLYAMINE AMINOHYDROLASE	Novel Protein sim. GBank gil4239787 emb CAA75437 - (Y15166) NADP-glutamate dehydrogenase [Pseudomonas aeruginosa]
80230771 (667, 668)	80057026 (669, 670)	80414319 (671, 672)	11090829 (673, 674)	95413134 (675, 676)	11398513 (677, 678)	80504149 (679, 680)	11075198 (681, 682)	80054196 (683, 684)	20466792 (685, 686)	80428870 (687, 688)	80258853 (689, 690)	79831058 (691, 692)
334	335	336	П	33.8	339	340	¥	342	343	84 44	345	346

265006, 265008, 265010, 265018, 263967, 263981	264602, 18108351, 18108387	265007	265009, 264769, 264689, 18108370	264769, 264905, 264908	264595	264604	264628	264909, 264595, 264683, 22279002	264909, 264591, 264592	264605	264768	264604, 264769	264594	22278996, 264259, 29331822, 29331824, 264605, 55811957, 265022	264688	26456R	264769, 264602, 264604, 264508, 264762, 264638, 264486
UNCLASSIFIED	transport		nuclease	glycoprotein	protease	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transcriptfactor	oxidase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transferase	amylase		dehydrogenase -
						Contains protein domain (PF00449) - UNCLASSIFIED Urease			Contains protein domain (PF00072) - transcriptfactor Response regulator receiver domain								Contains protein domain (PF00420) - dehydrogenase NADH-ubiquinone/plastoquinone oxidoreductase chain 4L
Novel Protein sim. GBank gl[731675 sp P38795 YHN4_YEAST - HYPOTHETICAL 80.7 KD PROTEIN IN ERG7-NMD2 INTERGENIC REGION	Novel Protein sim. GBank gij1073610 pir S47672 - ugpB protein - Escherichia coli		Novel Protein sim. GBank gij3261599 emb CAB00917 - (Z77137) hypothetical protein Rv1277 [Mycobacterium tuberculosis]	Novel Protein sim. GBank gil2959367 jemb CAA17921 - (AL022117) hypothetical protein (Schizosaccharomyces pombe)	Novel Protein sim. GBank gi[4416302]gb[AAD20307] - [AF105716] copia-type pol polyprotein [Zea mays]				Novel Protein sim. GBank gil115157 sp P16574 BVGA_BORPE - VIRULENCE FACTORS PUTATIVE POSITIVE TRANSCRIPTION REGULATOR BVGA	Novel Protein sim. GBank gil497637 (J03939) - cytochrome oxidase d subunit I [Escherichia coil]	Novel Protein sim. GBank gi 2290990 (AF006000) - Brg1 Bordetella pertussis]			Novel Protein sim. GBank gij3510639 (AF049344) - UDP- GalNAc:polypeptide N-acetylgalactosaminyltransferase T5 IRattus norveoicus	Novel Protein sim. GBank gij113764 sp P25718 AMY1_ECOLI - ALPHA-AMYLASE PRECURSOR (1.4-ALPHA-D-GLUCAN GLUCANOHYDROLASE)		Novel Protein sim. GBank gi[2829816 sp[P95171 NUOK_MYCTU - NADH DEHYDROGENASE I CHAIN K (NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 11) (NUO11)
79158195 (693, 694)	80020208 (895, 696)	17282112 (697, 698)	80502370 (699, 700)	80501805 (701, 702)	11611585 (703, 704)	80061653 (705, 706)	56626130 (707, 708)	80046344 (709, 710)	80043835 (711, 712)			80501488 (717, 718)	80026748 (719, 720)	-	ļ		82443593 (727, 728)
347	9	848		351	352		П	355				٦	9			363	

365	88040288 (729, 730)	Novel Protein sim. GBank gil4929268 gb AAD33924.1 -	Contains protein domain (PF00412) -		264488, 21906766, 21906767, 55811576,
		(AF144237) LOMP protein [Homo sapiens]	LIM domain containing proteins		21906769, 29148629, 22278995, 22278996,
					265020, 265022, 264634, 264691, 264593,
					33657023, 33657402, 264693, 264639,
					264594, 29331824, 264758, 18108385,
					29331827, 87168559, 265018, 22279000,
					265019, 264482, 264761, 264681, 18108351
366	81821838 (731, 732)				265017, 264757
367	95357471 (733, 734)		Contains protein domain (PF01602) - glycoprotein	glycoprotein	60424179, 65274572, 56182575, 22278994,
		gi 4503843 ref NP_003908.1 pG2AD - UNKNOWN	Adaptin N terminal region		56994075, 22278998, 264259, 29331822,
					29331824, 56182181, 60424269, 66714117.
					29331825, 60432289, 29331826, 29331827,
					29331828, 264905, 264828, 56182435,
					265006, 264512, 265008, 264591, 55812038,
			-		55811386, 265010, 87168559, 265017.
					265018, 264604, 265019, 55811150, 264448,
_					264369, 264288, 264686, 264768, 56181562,
					21906768, 21906769, 55811957, 35695917,
					265022, 60170615, 33657023, 65274620,
					18108365, 263967, 33657109, 33657349.
					35695763, 264628, 18108376, 55811576,
					65274791, 35695855, 56182323, 83373044,
					60432113, 264563, 264564, 264567
368	79607265 (735, 736)				264509
369	95292917 (737, 738)	Novel Protein sim. GBank		UNCLASSIFIED	264508, 264604, 264605, 264636
		gij3913029jspjP94967JALR_MYCSM - ALANINE	•	-	
		RACEMASE			
370	88090966 (739, 740)	Novel Protein sim. GBank gij3249559 (AF018261) - EH			264905, 264592, 264605, 264766, 264691
	٠	domain binding protein Epsin [Rattus norvegicus]			ſ
371	95292599 (741, 742)	Novel Protein sim. GBank gij2995299 emb CAA18328 -	Contains protein domain (PF01715) - Iransferase	transferase	264905, 264906, 264510, 264600, 264601,
		(AL022268) putative tRNA delta(2)-	IPP transferase		264602, 264603, 265018, 264604, 264605,
		isopentenylpyrophosphate transferase (Streptomyces			265021, 264692, 264636, 264564
		coelicolor)			
372	80021107 (743, 744)	Novel Protein sim. GBank			264564
		gi 2506393 sp P31576 FIXX_ECOLI - FERREDOXIN LIKE PROTEIN			
373	79863766 (745, 746)				264909
374	79847568 (747, 748)	Novel Protein sim. GBank gij3341640 emb CAA13164 -		UNCLASSIFIED	264905, 264906
		(AJ231122) z61f [Vibrio cholerae]			
375	91230181 (749, 750)	Novel Protein sim. GBank gi[5456934 gb AAD43716.1] -		cadherin	65274572, 264259, 29331826, 56182435,
		(AF152322) protocadherin gamma A2 [Homo sapiens]			60433356, 60433438, 264757, 55812038, 264758, 55811957, 264690, 33657023
376	80505214 (751, 752)	Novel Protein sim. GBank gil 1805408idbil BAA08970l -		UNCLASSIFIED	264769
	•	(D50453) homologues to nitrile hydratase region 3'-			
		hypothetical protein P47K of P. chlororaphis [Bacillus			
	170000000000000000000000000000000000000	SUDDING			20000
377	10339083 (753, 754)				Z64806

				l	
9/2	(dc), cc) (cc) (dc)			Olrico Olrico	203008, 204333
3.78 B	(40503437 (757, 758)	Novel Protein sim. GBank gij10/6013 pir A49930 - carb	Contains protein domain (PF00289) - Synthase		264769
		protein homolog - Mycobacterium bovis (strain BCG) (fragment)	Carbamoyi-phosphate synthase (CPSase)		
380	80060937 (759, 760)	Novel Protein sim. GBank gil216556 dbj BAA02174 - (D12651) glucose dehydrogenase [Escherichia coli]	Contains protein domain (PF01011) - dehydrogenase PQQ enzyme repeat		264604
381	11769027 (761, 762)			UNCLASSIFIED	264684
382	80054377 (763, 764)				264592
383	83259025 (765, 766)	Novel Protein sim. GBank gil3327136 db BAA31636 - (AB014561) KIAA0661 protein [Homo sapiens]			264595, 265017, 265021, 264638, 87168518, 22279002
384	95314255 (767, 768)			UNCLASSIFIED	264259, 29331822, 60432289, 29331827, 264288, 264766, 263967, 65274791,
					35695855, 263981, 83373044, 264567
385	10237679 (769, 770)				264692
386	79633434 (771, 772)	Novel Protein sim. GBank gi 1073456 pir S47810 - probable alcohol dehydrogenase (EC 1.1.1.1) - Escherichia coli	1073456 pir S47810 - probable Contains protein domain (PF00465) - dehydrogenase 1.1.1.1 - Escherichia coli Iron-containing alcohol dehydrogenases		264906
387	17960637 (773, 774)	Novel Protein sim. GBank gi[1460074 emb CAB01049] - (Z77250) hypothetical protein Rv2566 [Mycobacterium	Contains protein domain (PF01841) - UNCLASSIFIED Transglutaminase-like superfamily		264760
١	() the section of the	(uperculosis)	1000 D		25505385 264005 66712503 60432330
8	6//413/6 (//3, //6)	Nover Protein sim. Gbank gil4240169[db] BAA/4663.1 - (AB020647) KIAA0840 protein [Homo sapiens]	Contains protein domain (PF00040) - nomeobox F-box domain.	пошеорох	33030200, 204303, 007 12302, 00432223, 264593, 60433356, 264686, 264688,
					21906765, 264691, 22279000, 264482
388	79316971 (777, 778)			UNCLASSIFIED	18108394, 22278996, 264630, 264556, 22279002
390	80079949 (779, 780)			UNCLASSIFIED	264600
381	7657302 (781, 782)	Novel Protein sim. GBank gi 854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]			264482
392	79796056 (783, 784)			UNCLASSIFIED	264908
393	33206031 (785, 786)	Novel Protein sim. GBank gi[3378523]emb CAA08867 - (AJ009832) cyclomattodextrinase glucanotransferase Thermotoga neapolitana		synthase	264602, 21906764
394	10104463 (787, 788)				264693
392	80229010 (789, 790)			UNCLASSIFIED	264508, 264563
396	20436224 (791, 792)	Novel Protein sim. GBank gi 2677780 (U70327) - unknown [Paretroplus polyactis]	Contains protein domain (PF00047) - struct Immunoglobulin domain	struct	264556
397	80417014 (793, 794)	Novel Protein sim. GBank gil4507909 ref NP_000368.1 pVVAS - VViskott-Aldrich syndrome (ecezema-thrombocytopenia)			265007, 265009, 264508, 264556, 264629, 264766
398	91230517 (795, 796)	Novel Protein sim. GBank gij1518458 (U45998) - mitochondrial solute carrier [Onchocerca volvulus]	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	transport	18108398, 22278995, 22278996, 56994075, 22278999, 264259, 29331824, 29331826, 264905, 264908, 265007, 265008, 265009, 21906754, 33657084, 265017, 264448, 264288, 264766, 21906765, 21906766, 21906767, 265020, 265021, 33657023, 33657109, 264628, 35696423, 35695655, 264952, 18108380, 264567, 18108391

264592, 264595	18108394, 56182575, 22278995, 22278997, 22278999, 264259, 29331824, 265006, 265007, 265009, 60432229, 33657402, 21908754, 265010, 265017, 265018, 265019, 18108351, 18108387, 21906765, 265021, 265022, 264691, 284634, 264636, 60170384, 66182227	264594	65274572, 56182575, 35696286, 22278996, 22278998, 264093, 264259, 29331822, 29331824, 29331824, 29331824, 29331824, 29331825, 29331828, 60432289, 29331827, 29331826, 264907, 264909, 265006, 264511, 265007, 265008, 264910, 264591, 33657402, 80433356, 60433438, 264596, 21906754, 52644296, 265019, 256011, 265018, 264288, 264684, 264766, 265017, 2896769, 21906765, 21906766, 21906767, 21906768, 21906765, 21906766, 21906767, 21906768, 21906769, 55811957, 265020, 265021, 265022, 264690, 265021, 265022, 264690, 265021, 265022, 264590, 18108370, 264633, 18108376, 264639, 22279002, 264563, 264566, 24566, 24266, 244668, 264639, 18108385, 264659, 264659, 264659, 264659, 264659, 264665, 264669, 264668, 244666, 244	264768, 264632, 264639, 264563	265009, 264682	18108357, 264693	264769	264600	264259
Kinase	- cadherin	transport	phosphatase	UNCLASSIFIED	polymerase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	helicase
Contains protein domain (PF00370) - kinase FGGY family of carbohydrate kinases	Contains protein domain (PF00560) - cadherin Leucine Rich Repeat		Contains protein domain (PF00017) - phosphatase Src homology domain 2			Contains protein domain (PF00159) - UNCLASSIFIED Pancreatic hormone peptides			
Novel Protein sim. GBank gij3358091 jdbj BAA31995 - (AB015974) glycerol kinase [Pseudomonas tolaasii]	Novel Protein sim. GBank gij728335 sp p39192]ALU5_HUMAN - !!!! ALU SUBFAMILY Leucine Rich Repeat SC WARNING ENTRY !!!!	Novel Protein sim. GBank gil49282921gbJAAD33527.1JAF13211 - (AF132117) FhuA [Staphylococcus aureus]	Novel Protein sim. GBank gij5689487 dbj BAA83027.1 - (AB028998) K!AA1075 protein [Homo sapiens]		Novel Protein sim. GBank gi 2661649 emb CAA15755 - (AL009198) dnaE2 [Mycobacterium (uberculosis]				Novel Protein sim. GBank gi 3047117 (AF058919) - similar to ATP-dependent RNA helicases [Arabidopsis thaliana]
80055278 (797, 798)	94117490 (799, 800)			80439913 (805, 806)	11809865 (807, 808) 79471280 (809, 810)	79634172 (811, 812)	80478229 (813, 814)	Ĭ	5640527 (817, 818)
388	6 6	- - -	402 2	4 03	404		407	8	6 0

Ş	1069 640 640 6201	Married Desde Sim Contact Characterists & Acceptant			
	00001430 (014, 900)	(D29801) Unknown [Mus musculus]			26469, 3264630, 3264684, 30101086, 35696286, 32645080, 29331822, 29331824, 35696286, 52645080, 29331822, 29331824, 265970, 264508, 264509, 264905, 264905, 264907, 264907, 264908, 2644045, 264909, 264900, 265001, 265001, 265008, 264910, 33657402, 264768, 264768, 264910, 33657402, 264762, 264768, 264687, 264768, 264768, 264768, 264769, 3365702, 265020, 264030, 264639, 264629, 264638, 264639, 264639, 264639, 264639, 264638, 264638, 264639, 264636, 264486
=	80501670 (821, 822)			UNCLASSIFIED	
412	80241662 (823, 824)				264907, 264910, 263973, 22279002
413	11076446 (825, 826)	Novel Protein sim. GBank gij3261784 emb CAB08997 - (29558) htpX [Mycobacterium tuberculosis]		ydə	264605
414	82050554 (827, 828)	Novel Protein sim. GBank		dehydrogenase	18108374 264760 264769 264602 264638
		gi 129036 sp P20707 ODO1_AZOVI - 2-OXOGLUTARATE DEHYDROGENASE E1 COMPONENT (ALPHA- KETOGLUTARATE DEHYDROGENASE)			264603, 264909, 264605
<u>4.</u>	84453144 (829, 830)	Novel Protein sim. GBank gil4868350[gb]AAD31273.1 AF13202 - (AF132025) rhophilin		UNCLASSIFIED	264908, 87168518
		[Drosophila melanogaster]			
416	80402775 (831, 832)	Novel Protein sim. GBank gi[2555172 (AF025543) - ArcC, carbamate kinase IRhizobium etti]		kinase	264488, 264600, 264602, 264764, 264636
417	20153797 (833, 834)	Novel Protein sim. GBank	Contains profein domain (PE00145) -		264605
		gilt709171spl952311MTX2_XANOR - MODIFICATION METHYLASE XORII (CYTOSINE-SPECIFIC METHYLTRANSFERASE XORII) (M.XORII)			CORPOR
418	94125841 (835, 836)			UNCLASSIFIED	264689, 264693
419	95314273 (837, 838)			collagen	264908, 264910, 264764, 264639
420	37036349 (839, 840)		Contains protein domain (PF00300) - phosphatase Phosphoglycerate mutase family	phosphatase	264769
421	95292942 (841, 842)	Novel Protein sim. GBank gil2916942 emb CAA17580 - (AL021999) hypothetical protein Rv0981 [Mycobacterium tuberculosis]	Contains protein domain (PF00072) - Response regulator receiver domain	phosphatase	264906, 264600, 264601, 264603, 264604, 264760, 264769
422	79471293 (843, 844)	Novel Protein sim. GBank GP600 GP1221752!sp 000767 CH61_STRAL - 60 KD CHAPERONIN TCP-1/cpn60 chaperonin family (PROTEIN CPN60 1) (GROEL PROTEIN 1) (HSP58)	Contains protein domain (PF00118) - eph TCP-1/cpn60 chaperonin family	ф	22278996, 264682, 18108376, 18108387
423	79604948 (845, 846)			UNCLASSIFIED	264509

3	1010 5101 53333005				
	(940, 940)	Novel Protein sim. Coank gij4826814[ref]NP_004977.1[pKTN1 - kinectin 1 (kinesin receptor)		struct	265019
425	80431450 (849, 850)	Novel Protein sim. GBank gi 1703701 bbs 178462 - KRP5=kinesin-related protein [rats, testes, Peptide Partial, 167 aa]	Contains protein domain (PF00225) - struct Kinesin motor domain		264909, 265007, 55811386, 264768. 55810764
428	80064522 (851, 852)				264605, 264559
427	80057232 (853, 854)	Novel Protein sim. GBank gi 231829 sp P29929 COBN_PSEDE - COBN PROTEIN		UNCLASSIFIED	264603, 264636
428	79487798 (855, 856)			UNCLASSIFIED	264683
429	80091252 (857, 858)	Novel Protein sim. GBank gil81286 pir S22697 - extensin - Volvox carteri (fragment)			35696423, 35695763, 35695855, 265017. 264564, 264762
430	80504192 (859, 860)	Novel Protein sim. GBank gij1806154 jemb CAB06451 -		reductase	264508, 264905, 264509, 264908, 264909
		(284395) hypothetical protein Rv0688 [Mycobacterlum			265008, 264600, 264687, 264769, 264689,
431	20624249 (861, 862)				264656
432	16525372 (863, 864)				255020
433	81494303 (865, 866)	Novel Protein sim. GBank gij3123552 emblCAA18609 -		UNCLASSIFIED	264907 264908 264909 264910 264592
		(AL022578) dJ393P12.2 (hypothetical Proline-rich protein KIAA0269 LIKE) IHomo sapiensl			264595, 264758, 264604, 264760, 264762, 264763
434	94326323 (867 868)	Novel Protein sim Chank	Contains acotoin domain (DE00160)	T	ECO1000 CO10010 CO10010 CO1000
_	(222,122)	gil2495272lsplQ99626lCDX2 HUMAN - HOMEOBOX	Contains protein comain (Probles) - CNCLASSIFIED PH domain		35696052 55810764 55811576 65274791
		PROTEIN CDX-2 (CAUDAL-TYPE HOMEOBOX PROTEIN			35695855, 60432113, 55811150, 264636.
		2) (CDX-3)			264766
435	80502738 (869, 870)	Novel Protein sim. GBank		transport	264595, 264769
		gij114105jspjP08532jaRah_ECOLI - L-ARABINOSE TRANSPORT SYSTEM PERMEASE PROTEIN ARAH			
438	41085953 (871, 872)			UNCLASSIFIED	265020, 22279002
437	11399291 (873, 874)				264593
438	11773835 (875, 876)				264686
439	80019495 (877, 878)	Novel Protein sim. GBank gil3242702 (AC003040) -			264905 264600 264602 264604
		hypothetical protein [Arabidopsis thaliana]			101-300, 101-001, 101-001, 101-001
0	79841062 (879, 880)	Novel Protein sim. GBank gi 2291232 gb AAB65351.1 -	Contains protein domain (PF00004) -	ATPase_associated	Contains protein domain (PF00004) - ATPase_associated 35696052, 264905, 264908, 264909, 265011,
		(AF016427) Contains similarity to Pfam domain: PF00004 ATPases associated wi (AAA), Score=268.1, E-value=3.7e-77, N=1 [Caenorhabdiiis [cellular activities (AAA)	ATPases associated with various cellular activities (AAA)		35698423
		elegans]			
<u>\$</u>	20396935 (881, 882)	Novel Protein sim. GBank			264605
		gripopagagajiAAD45904.1 AF 15132 - (AF 151328) nistidine kinase CstS (Corynebacterium diphtheriae)			
445	85281058 (883, 884)	Novel Protein sim. GBank gi 1184790 (U46068) - von Ebner		UNCLASSIFIED	29331830, 264909
		minor salivary gland protein [Mus musculus]			
4	82456427 (885, 886)	Novel Protein sim. GBank gij5689893 emb CAB52056.1 -		UNCLASSIFIED	35696052, 264508, 264906, 264512, 264604.
		(AL109732) putative ATP-binding RNA helicase [Streptomyces coelicolor A3(2)]			264762, 264769, 264689, 264636
4 4 4	11395897 (887, 888)	Novel Protein sim. GBank gij1783249 dbj BAA11726 -		UNCLASSIFIED	264591
		(O83026) homologous to citrate-sodium symport (citrate transporters): hypothetical (Bacillus subtilis)		•	
				1	

445	79552709 (889, 890)			UNCLASSIFIED	264693
446	79810937 (891, 892)	Novel Protein sim. GBank gi[5531272 emb CAB50897.1 - (AJ243800) WSC4 homologue [Kluyveromyces lactis]			264509
447	80438888 (893, 894)	Novel Protein sim. GBank gi 538413 (L36315) - zinc finger	Contains protein domain (PF00096) - transcriptfactor	transcriptfactor	264768, 55811576
448	80238110 (895, 896)	Process (International State of State o	Contain protein domain (PF00551) - dehydrogenase Formy transferase	dehydrogenase	264508, 264600, 264603, 264605, 264682, 264769, 18108362, 264634, 18108387
449	20460634 (897, 898)	Novel Protein sim. GBank gij118794jspjP10443jDP3A_ECOLi - DNA POLYMERASE III ALPHA CHAIN		polymerase	264605, 264559
450	84631210 (899, 900)	Novel Protein sim. GBank gil4589506 dbj BAA76775.1 - (AB023148) KIAA0931 protein [Homo sapiens]	Contains protein domain (PF00481) - phosphatase Protein phosphatase 2C	phosphatase	65274572, 22278998, 29331824, 29331826, 264906, 264910, 264592, 52646317, 265017, 21906767, 55811957, 56578486, 22279002
155	21433609 (901, 902)			UNCLASSIFIED	264486
452	10267276 (903, 904)				264692
453	52560096 (905, 906)	Novel Protein sim. GBank gi/2650614 (AE001104) - conserved hypothetical protein [Archaeoglobus fulgidus]		UNCLASSIFIED	264907, 264600
454	39523922 (907, 908)	Novel Protein sim. GBank gi[2493000]sp Q09450 SCOT_CAEEL - PROBABLE SUCCINYL-COA:3-KETOACID-COENZYME A TRANSFERASE PRECURSOR (3-OXOACID COA-TRANSFERASE)		transferase	264603
455	13089692 (909, 910)			UNCLASSIFIED	264687
458	79563081 (911, 912)			UNCLASSIFIED	264691
457	79831273 (913, 914)	Novel Protein sim. GBank gil4468699 emb CAB38153.1 - (AL035591) putative integral membrane export protein [Streptomyces coelicolor]			264905
458	79581227 (915, 916)	Novel Protein sim. GBank gij3411053 (AF034863) - synaptic scaffolding molecule [Rattus norvegicus]	Contains protein domain (PF00595) - kinase PDZ domain (Also known as DHR or GLGF).	kinase	55812038, 265010, 265018, 264681
459	80567359 (917, 918)	Novel Protein sim. GBank gi 4506075 ref NP_002733.1 pPRKC - protein kinase C, mu	Contains protein domain (PF00130) - Phorbol esters/diacylglycerol binding domain (C1 domain)	kinase	22278997, 264259, 29331826, 265018. 264448, 264369, 21906765, 35696423
460	79245890 (919, 920)	Novel Protein sim. GBank gij113158 sp P25516 ACO1_ECOLI - ACONITATE HYDRATASE 1 (CITRATE HYDRO-LYASE 1) (ACONITASE 1)		UNCLASSIFIED	264906
461	95287618 (921, 922)	Novel Protein sim. GBank gif168574lspIP42464 ATPB_CORGL - ATP SYNTHASE BETA CHAIN		synthase	264602, 264605, 264768, 264769, 265021, 33657023, 264559

264907	264905, 264906, 264908, 264909, 264910. 264591, 264595, 265011, 264632, 284635, 264636, 264637, 264638, 264639	264634	265017, 21906764, 265020	264605, 264559	264764	264692	264488	264602, 264769	265019	264596, 264685, 264557	264369	22278997, 264692, 264288	264907, 264908, 264511, 265009, 264762, 264448, 264636, 264638	65274572, 60432049, 264259, 264508,	52644045, 55812038, 264758, 255011, 264288, 264686, 52644229, 65274791, 264638, 264568	264636	264690, 264693	264686, 35695855, 265008, 264631, 264910, 264632, 264638, 265018, 264369, 264909	264369	264693	264909, 264686, 264768, 264693, 55811576, 56182323, 18108385
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	cathepsin		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	synthase	transport		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	glycoprotein		UNCLASSIFIED	UNCLASSIFIED	collagen	mapolymerase	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00391) - UNCLASSIFIED PEP-utilizing enzymes			Contains protein domain (PF00648) - cathepsin Calpain family cysteine protease	Contains protein domain (PF00417) - Ribosomal protein S3, N-terminal domain.										Contains protein domain (PF00560) - glycoprotein	Leucine Rich Repeat						
HPR	Novel Protein sim. GBank gij854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]		Novel Protein sim. GBank gi[5689776 emb CAB52137.1 - (AJ242832) calpain [Homo sapiens]	'5 emb CAB06470 - :rculosis]	Novel Protein sim. GBank gij548705 sp P36949 RBSB_BACSU - D-RIBOSE-BINDING PROTEIN PRECURSOR		Novel Protein sim. GBank gil2114024[emb CAB08957] - (295558) grcC1 [Mycobacterium tuberculosis]	Novel Protein sim. GBank gil2909459 emb CAA17347 - (AL021929) cobQ (Mycobacterium tuberculosis)	Novel Protein sim. GBank gij114921 sp P17447 BETT_ECOLI - HIGH-AFFINITY CHOLINE TRANSPORT PROTEIN		Novel Protein sim. GBank gil862343 (L10908) - Gcap1 gene product (Mus musculus)			Novel Protein sim. GBank	gi[5453656]ref[NP_006329.1 pGAC1 - glioma amplified on chromosome 1 protein (leucine-rich)				Novel Protein sim. GBank gi[1127551 (U18939) - orf2		Novel Protein sim. GBank gil4063042 (AF068065) - GP900; mucin-like glycoprotein [Cryptosporidium parvum]
79606589 (923, 924)	79796417 (925, 926)	82340151 (927, 928)	83005730 (929, 930)	20460645 (931, 932)	80409035 (933, 934)	52562208 (935 936)	19520527 (937, 938)	80502756 (939, 940)	17937351 (941, 942)	80047458 (943, 944)	20558793 (945, 946)	80593365 (947, 948)	82454665 (949, 950)	94143857 (951, 952)		70475023 (063 064)	70622483 (933, 934)	80189746 (957, 958)	79390729 (959, 960)	70624578 (061 962)	83050611 (963, 964)
462	463	Т	465	466	467	Т	469	470	471	472	473	Г	475	476		,,,,) } }	479	480	187	482

483	20293306 (965, 966)	Novel Protein sim. GBank gi 2104303 emb CAB08632 - (295387) hypothetical protein Rv2610c [Mycobacterium InharralAcis]	Contains protein domain (PF00534) - Glycosyl transferases group 1		264600
48	11618046 (967, 968)	Novel Protein sim. GBank gij3450883 (AF083334) - fibroin Antheraea nemvi		UNCLASSIFIED	264594
85	80191234 (969, 970)			UNCLASSIFIED	264369, 21906765, 22279000, 22279002
486	80059042 (971, 972)	Novel Protein sim. GBank gil5042272 emb CAB44526.1 - (AL078618) nuoF. NADH dehydrogenase subunit Streptomyces coelicolor		dehydrogenase	264604
487	11813339 (973, 974)				264638
488	91222383 (975, 976)	Novel Protein sim. GBank gij5724778 gb AAC53522.2 - Contains protein (AF012273) mo-type GTPase-activating protein moGAPX-1 RhoGAP domain [Mus musculus]	Contains protein domain (PF00620) - RhoGAP domain		264686, 66714117, 264768, 18108385, 55811576, 265006, 265008, 265009, 265019, 22279002, 264259, 18108370, 264907, 264764, 56182323, 264288, 264693
489	10867710 (977, 978)	Novel Protein sim. GBank gij3882223 dbj BAA34471.1 - (AB018294) KIAA0751 protein [Homo sapiens]		kinase	264639
490	95361124 (979, 980)	Novel Protein sim. GBank gil82091 pir A25494 - hydroxyproline-rich glycoprotein - tomato (fragment)		collagen	22278996, 29331822, 29331828, 264107, 264909, 264110, 265009, 264592, 264593, 60433356, 264288, 264693, 263974, 263976, 20281071, 60432113
491	80496412 (981, 982)	Novel Prolein sim. GBank gij2894206jembjCAA17072 - (AL021840) hypothetical protein Rv3258c [Mycobacterium tuberculosis]		UNCLASSIFIED	264769
28	87421264 (983, 984)				264600
8	11692942 (985, 986)			UNCLASSIFIED	264638
484	87726604 (987, 988)	Novel Protein sim. GBank gij5262605 emb CAB45743.1 - (AL080150) hypothetical protein [Homo sapiens]		UNCLASSIFIED	264489, 35696286, 60432289, 29331828. 35696052, 264509, 264905, 264906, 264907, 264908, 264909, 264510, 264511, 265009.
					264910, 33657402, 264762, 264764, 264768, 264769, 264769, 284688, 21906765, 21906769, 35695917, 265020, 264693, 33657109, 264629, 35695855, 264634, 264638
495	80028599 (989, 990)	Novel Protein sim. GBank gi 2791517 emb CA416054 - (AL021246) hypothetical protein Rv2477c [Mycobacterium tuberculosis]	Contains protein domain (PF00005) - transport ABC transporter	transport	264602, 264682, 264638
496	78985624 (991, 992)	Novel Protein sim. GBank gij2302811pdb 1R69 - 434 Repressor (Amino-Terminal Domain) (R1-69)	Contains protein domain (PF01381) - Helix-tum-helix		264601, 265021
497	78949661 (993, 994)			oxidase	265006

498	88095488 (995, 996)	Novel Protein sim. GBank gi 1145789 (U41662) - neuroligin 2 [Rattus norvegicus]	1145789 (U41662) - neuroligin Contains protein domain (PF00135) - esterase Carboxylesterases	esterase	264259, 29331826, 35698052, 264508, 264509, 264509, 264905, 264906, 264907, 264908, 264909, 264510, 265009, 264910, 264591, 265010, 264591, 33657402, 264786, 265083, 264764, 264767, 264768, 264687, 264769,
					21906767, 33657023, 284693, 264628, 284629, 35696423, 264630, 264632, 264634, 264635, 264558, 264563, 264565, 264566, 264566, 264566, 264566, 264567
488	20438222 (997, 998)	Novel Protein sim. GBank gil97480[pir][S19739 - integral membrane protein - Rhodobacter capsulatus		UNCLASSIFIED	264605
200	11076810 (999, 1000)				264605
	13418034 (1001, 1002)	13418034 (1001, 1002) Novel Protein sim. GBank gil5708250[embJCAB52363.1] - (AL 109747) putative integral membrane protein [Streptomyces coelicolor A3(2)]		UNCLASSIFIED	264688
502	80021176 (1003, 1004)		Contains protein domain (PF00342) - isomerase Phosphoglucose isomerase	isomerase	22278996, 265011, 264602, 264605, 264635
	20264483 (1005, 1006)			UNCLASSIFIED	264564
504	10887321 (1007, 1008)			UNCLASSIFIED	264687
	95003068 (1009, 1010)			UNCLASSIFIED	264369
506	16454292 (1011, 1012)	16454292 (1011, 1012) Novel Protein sim. GBank gi 4033509 sp P02598 CALM_TETPY - CALMODULIN	Contains protein domain (PF00036) - struct EF hand	struct	265010
507	20451598 (1013, 1014)			UNCLASSIFIED	264604
		gi[2501069]sp[Q46127]SYW_CLOLO - TRYPTOPHANYL- TRNA SYNTHETASE (TRYPTOPHANTRNA LIGASE) (TRPRS)			
508	79841424 (1015, 1016)			UNCLASSIFIED	264908
		gi 466068 sp P34618 YO82_CAEEL - HYPOTHETICAL 33.8 KD PROTEIN ZK1236.2 IN CHROMOSOME III			
509	(11776386 (1017, 1018)				264638
510	83373465 (1019, 1020)			UNCLASSIFIED	264687, 264639
511	(16525578 (1021, 1022)				265007
512	20399484 (1023, 1024)			UNCLASSIFIED	264565
		gi[2497419]sp P55635 Y4RB_RHISN - PUTATIVE INTEGRASERECOMBINASE Y4RB			
513	79457404 (1025, 1026)	79457404 (1025, 1026) Novel Protein sim. GBank gij1276897 (U41809) - cyclin J	Contains protein domain (PF00134) - cyclin	cyclin	264683, 264689, 35696423, 264639
514	79813805 (1027 1028)	79813805 (1027 1028) Novel Protein sim GBank di 1184790 (146068) - von Ebner	_	INCLASSIFIED	29331830 264909
		minor salivary gland protein [Mus musculus]			
515	79462591 (1029, 1030)				22278999, 264690
_	9862020 (1031, 1032)	Novel Protein sim. GBank gi[2127400 pir S65770 -		amylase	264910
I			***************************************		

517	95292994 (1033, 1034)	95292994 (1033, 1034) Novel Protein sim. GBank gil2983605 (AE000725) - ribose 51		isomerase	265018 264606 264764 264766 264687
		phosphate isomerase B [Aquifex aeolicus]			264691, 264565
518	8491831 (1035, 1036)	Novel Protein sim. GBank gil854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264487
519	91677886 (1037, 1038) 70860188 (1030, 1040)	Novel Protein sim. GBank gij5689365[dbj]BAA83073.1 - (AB024075) B120 [Homo sapiens]	Contains protein domain (PF01388) - dna_rna_bind ARID DNA binding domain	dna_ma_bind	52644507, 22278997, 22278998, 60432049, 264259, 52645080, 29331824, 66714117, 60424269, 29331826, 35696052, 264905, 29331830, 66712502, 264511, 265007, 264591, 60432229, 33657402, 60433438, 21906754, 33109954, 52644296, 8716859, 265017, 265018, 264604, 265019, 264681, 264448, 26439, 264298, 264685, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21806365, 21806365, 21806365, 21806365, 2564332, 565012, 33657023, 264632, 2486264, 18108335, 56526488, 60432113
3	79659166 (1039, 1040)				264769
521	11076821 (1041, 1042)	11076821 (1041, 1042) Novel Protein sim. GBank gi 1169126 sp P46839 CTPA_MYCLE - CATION- TRANSPORTING P-TYPE ATPASE A		transport	264605
252	80435060 (1043, 1044)	Ų.	Contains protein domain (PF00294) - kinase pfkB family carbohydrate kinase	kinase	264905, 264768
523	18356013 (1045, 1046)			UNCLASSIFIED	264629
524	80261805 (1047, 1048)	80261805 (1047, 1048) Novel Protein sim. GBank gil4033608 dbj BAA35136 - (AB012308) B2HC [Anthocidaris crassispina]		ATPase_associated	ATPase_associated 264092, 264596, 265011
525	79610046 (1049, 1050)				264907
256	36827630 (1051, 1052) 	36827630 (1051, 1052) Novel Protein sim. GBank gil4106610 emb CAA21365 - (AL031866) ORF42, len=386 aa , similarity to an aminotransferase, in P95957 Sulfolobus solfataricus, (401 aa), 33.1% identity in 393 aa overlap, Fasta scores: opt:468, E(): 8.5e-24, in Q64602 R. norvegicus, (425 aa), 28.6% ident		UNCLASSIFIED	264758
527	80504729 (1053, 1054)			UNCLASSIFIED	264769
528	65484134 (1055, 1056)				56182575, 265017, 265018
529	17936810 (1057, 1058)	17936810 (1057, 1058) Novel Protein sim. GBank gij731088jspjP24215jUXUA_ECOLI - MANNONATE DEHYDRATASE (D-MANNONATE HYDROLASE)			265019
230	10887336 (1059, 1060)	Novel Protein sim. GBank gil42144 emb CAA25200 - (X00513) NusA protein (nusA) [Escherichia coli]		UNCLASSIFIED	264687
531	80226576 (1061, 1062)			Γ	264555, 264556, 264557, 264558, 18108385
532	90933444 (1063, 1064)	90933444 (1063, 1064) Novel Protein sim. GBank gi[5262640]emb[CAB45758.1] - [CAL080170) hypothetical protein [Homo sapiens]	Contains protein domain (PF00622) - UNCLASSIFIED SPRY domain		264488, 264490, 264259, 264592, 264760, 265021, 264690, 263976, 264558

533	87761531 (1065, 1066)	87761531 (1085, 1066) Novel Protein sim. GBank glj4883636jgbjAAD31593.1jAF11229 - (AF112299) integral inner nuclear membrane protein MAN1 (Homo capiens)			264907, 264909, 264768, 35695917, 264630, 264555
534	82368264 (1067, 1068)	82368284 (1087, 1068) Novel Protein sim. GBank gi[2995332 emb CAA04606.1 - (AJ001206) pep1 [Streptomyces coelicotor]		UNCLASSIFIED	264905, 265011, 264601, 264602, 264605, 264762, 264768, 265020, 264693, 264636
535	79641850 (1069, 1070)	79641850 (1069, 1070) Novel Protein sim. GBank gij3878636jembjCAA88953j - (Z49128) similar to cAMP-dependant protein kinase; cDNA EST EMBL:100719 comes from this gene; cDNA EST yk46548.3 comes from this gene; cDNA EST yk46548.5 comes from this gene; cDNA EST yk49214.3 comes from this gene; cDNA EST yk49214.3 comes from this gene; cDNA EST yk49214.3 comes from	Contains protein domain (PF00069) - ATPase_associated 264906 Eukaryotic protein kinase domain	ATPase_associated	264906
536	79907207 (1071, 1072)	79907207 (1071, 1072) Novel Protein sim. GBank gi[2495628]sp P55757 YOHL_SERMA - HYPOTHETICAL 10.1 KD PROTEIN IN BIOA 5'REGION		reductase	18108376, 264905, 264906, 264907, 264909
537	94147448 (1073, 1074)				265008, 264605, 65274791
538	87821963 (1075, 1076)	87821963 (1075, 1076) Novel Protein sim. GBank gill 34920 spjP21997 SSGP_VOLCA - SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185)	Contains protein domain (PF00595) - collagen PDZ domain (Also known as DHR or GLGF).		29331822, 29331824, 29331825, 29331826, 29331827, 264908, 52644045, 33657402, 265017, 264762, 264683, 264288, 264685, 21906765, 35695763, 264558, 60170394, 264559, 22279002
539	28396269 (1077, 1078)	28396269 (1077, 1078) Novel Protein sim. GBank gi 2498433 sp Q12341 HAT1_YEAST - HISTONE ACETYLTRANSFERASE		histone	264602, 265019
540	79637077 (1079, 1080)				264693
541	87762268 (1081, 1082)	87762268 (1081, 1082) Novel Protein sim. GBank gij3882241 dbj BAA34480.1 - (AB018303) KIAA0760 protein [Homo sapiens]	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type		18108394, 22278997, 22278998, 264259, 264112, 265009, 33657402, 55812038, 52646317, 265017, 21906765, 264693, 55811576, 264635, 56526486, <u>264566</u>
542	95295836 (1083, 1084)	95295836 (1083, 1084) Novel Protein sim. GBank gij5042272 emb CAB44526.1 - (AL078618) nuoF, NADH dehydrogenase subunit [Streptomyces coelicolor]			264910, 265018, 264689, 264638, 264486
543	79796290 (1085, 1086)				264602, 264908
544	20437191 (1087, 1088)	20437191 (1087, 1088) Novel Protein sim. GBank gi 2791398 emb CAA15994 - (AL021184) hypothetical protein Rv1464 [Mycobacterium tuberculosis		UNCLASSIFIED	264605
545	80434504 (1089, 1090)				264768, 264634, 264907, 264592, 264909
546	80249016 (1091, 1092)	80249016 (1091, 1092) Novel Protein sim. GBank gil4887211[gblAAD32237.1 AF14744 - (AF147449) penicillin binding protein 1B [Pseudomonas aeruginosa]		,	264600, 264602, 21906765
547	11077563 (1093, 1094)	11077563 (1093, 1094) Novel Protein sim. GBank gij1350855 sp P19176 RPOC_PSEPU - DNA-DIRECTED RNA POLYMERASE BETA' CHAIN (TRANSCRIPTASE BETA' CHAIN) (RNA POLYMERASE BETA' SUBUNIT)		глароlутегаѕе	264604
548	82114936 (1095, 1096)	82114936 (1095, 1096) Novel Protein sim. GBank gil2330021 (AF019250) - kinesin- related protein; KRP; Costal2 [Drosophila melanogaster]		UNCLASSIFIED	264488, 264905, 264910, 264760, 264693, 264639, 264563, 264564

22278996, 22278997, 22278998, 22278999, 22278996, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22228999, 22228999, 22228999, 22228999, 2222891, 2222899, 2222891, 222289, 222892, 2228, 22892, 222892, 222892, 2228, 2228, 2228, 2228, 2228, 2228, 2228, 22	264688		ED 264689, 264639, 264563		264906	264762	or 264508, 264605, 264559	se 264488	ED 264602	ED 264634
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	glycoprotein	kinase	transport	transcriptfactor	dehydrogenase	UNCLASSIFIED	UNCLASSIFIED
				Contains protein domain (PF00008) - glycoprotein EGF-like domain		Contains protein domain (PF00358) - transport phosphoenolpyruvate-dependent sugar phosphotransferase system, EIIA 1				
95421904 (1097, 1098) Novel Protein sim. GBank gil4337460[gb AAD18133] - (AF056195) neuroblastoma-amplified protein [Homo sapiens]		80439990 (1101, 1102) Novel Protein sim. GBank gi 3122893 sp P94985 SYFB_MYCTU - PHENYLALANYL- TRNA SYNTHETASE BETA CHAIN (PHENYLALANINE TRNA LIGASE BETA CHAIN) (PHERS)		80106002 (1105, 1106) Novel Protein sim. GBank gi 552087 (M33753) - crumbs protein [Drosophila melanogaster]	79618379 (1107, 1108) Novel Protein sim. GBank gi 5019771 gb AAD37857.1 AF13326 - (AF133263) histidine protein kinase-response regulator hybrid protein CvgSY Pseudomonas syringae pv. syringae]	Novel Prolein sim. GBank gif131515 sp P02908 PTGA_SALTY - PTS SYSTEM, GLUCOSE-SPECIFIC IIA COMPONENT (EIIA-GLC) (GLUCOSE-PERMEASE IIA COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, A COMPONENT)	20457127 (1111, 1112) Novel Protein sim. GBank gij3914014 sp P96380 MFD_MYCTU - TRANSCRIPTION- REPAIR COUPLING FACTOR (TRCF)	19523405 (1113, 1114) Novel Protein sim. GBank gil5042273 emb CAB44527.1 - (AL078618) nuoE, NADH dehydrogenase subunit [Streptomyces coelicolor]	20724429 (1115, 1116) Novel Protein sim. GBank gij1170933 sp P45331 METE_HAEIN - 5- METHYLTETRAHYDROPTEROYLTRIGLUTAMATE HOMOCYSTEINE METHYLTRANSFERASE (METHIONINE SYNTHASE, VITAMIN-B12 INDEPENDENT ISOZYME) (COBALAMIN-INDEPENDENT METHIONINE SYNTHASE)	80084353 (1117, 1118) Novel Protein sim. GBank gil4980567Igb AAD35173.1 AE00169 - (AE001694) iron(III) ABC transporter, permease protein [Thermotoga maritima]
95421904 (1097, 1098)	10886616 (1099, 1100)	80439990 (1101, 1102)	94672870 (1103, 1104)	80106002 (1105, 1106)	79618379 (1107, 1108)	78996347 (1109, 1110)	20457127 (1111, 1112)	19523405 (1113, 1114)	20724429 (1115, 1116)	80084353 (1117, 1118)
82 8	220	551	552	223	55 4	555	556	557		559

090	80059417 (1159, 1160)				22278999, 35696052, 264555, 264556, 264558
581	79230833 (1161, 1162	79230833 (1161, 1162)		UNCLASSIFIED	265008, 264564
285	80049617 (1163, 1164	Novel Protein sim. GBank gi 3243131 (AF045777) - titin Drosophila melanogaster	Contains protein domain (PF00047) - struct Immunoglobulin domain	struct	265021, 264555, 264557
583	79321392 (1165, 1166		 	transport	264594
584	79845024 (1167, 1168)			UNCLASSIFIED	264488, 264906, 264766, 264687, 35696423
585	79581454 (1169, 1170	79581454 (1169, 1170) Novel Protein sim. GBank gij3882221/dbjjBAA34470.11 - (AB018293) KIAA0750 protein [Homo sapiens]		UNCLASSIFIED	265018, 264684, 21906769
286	38277486 (1171, 1172)	(1)		UNCLASSIFIED	264908, 265007
287	80497359 (1173, 1174	80497359 (1173, 1174) Novel Protein sim. GBank gil4467250 emb CAB37575 -		hydrolase	264600, 264602, 264605, 264769, 264690.
		(AL035569) probable Glu-tRNA Gln amidotransferase subunit [Streptomyces coelicolor]			264557
288	79557239 (1175, 1176	79557239 (1175, 1176) Novel Protein sim. GBank gil5689519 dbj BAA83043.1 - (AB029014) KIAA1091 protein [Homo sapiens]		UNCLASSIFIED	265020, 264692
289	79805828 (1177, 1178)			UNCLASSIFIED	22278996, 264907, 264909, 264510, 265009, 265010, 264087, 264769, 35695917, 18108376, 264634, 264638, 264638
280	79815629 (1179, 1180)			INCI ASSIFIED	SEARCH SEARCH
591	10313540 (1181, 1182	10313540 (1181, 1182) Novel Protein sim. GBank gil2143293 emblCAB09390 -		mapolymerase	264691
		(295972) rpoB [Mycobacterium tuberculosis]			
292	13889767 (1183, 1184			MHC	263972
283	82348699 (1185, 1186	82348699 (1185, 1186) Novel Protein sim. GBank gij4511983jgbJAAD21543.1J -		dehydrogenase	264511, 264762, 264769, 264486
		(AF088896) electrotransfer ubiquinone oxidoreductase [Zymomonas mobilis]			
594	20212392 (1187, 1188	20212392 (1187, 1188) Novel Protein sim. GBank gi 1272368 (U51896) - LígE		UNCLASSIFIED	264605
	4000 4004 4400	(Vidno paranaemolyticus)			
0 6 	10064054 (1189, 1190 	10054054 (1159, 1190) Novel Prolein sim. GBank gi 131490 sp P20966 PTFB_ECOLI - PTS SYSTEM.			264769
		FRUCTOSE-SPECIFIC IIBC COMPONENT (EIBC-FRU) (FRUCTOSE-PERMEASE IIBC COMPONENT)			
296	13085170 (1191, 1192)	9)		UNCLASSIFIED	264636
597	80259003 (1193, 1194)	1		UNCLASSIFIED	264592
298	94140216 (1195, 1196)			UNCLASSIFIED	264758, 55810764, 264555, 264556, 264637,
599	20385137 (1197 1198	20385137 (1197, 1198) Novel Protein sim GBank		INC. ACCIEIED	264603
		gilt25;29(spiPo4951(kDS)_ECOLI - 3-DEOXY-MANNO-OCTULOSONATE CYTIDYLYLTRANSFERASE (CMP-KDO SYNTHETASE) (CMP-2-KETO-3-DEOXYOCTULOSONIC ACID SYNTHETASE) (CMP-2-KETO-3-DEOXYOCTULOSONIC			
000	10357663 (1199, 1200)	-			264906
601	79610404 (1201, 1202	79610404 (1201, 1202) Novel Protein sim. GBank gi[2127414 pir S60064 -		UNCLASSIFIED	264510
		hypothetical protein 2 - Corynebacterium glutamicum			

265007	264595	264758	264605	264764	264508, 264906, 85658542, 264682, 264687, 264689, 284534, 18108376, 35696423, 264636, 264638	264682	264605	264692	264508, 264905, 264907, 264508, 264909, 264511, 264910, 264558, 264564, 264684, 264766, 264689, 264628, 264635, 264636, 264637, 264636, 264637, 264588, 264637, 264588, 264637, 264588, 264637, 264588, 264637, 264588, 264638, 264688	264600, 264601, 264604, 264769, 264558,	264565	264369	18108372, 264563	264600, 264602, 264629	264605	52645156, 21906765, 35696423, 21906768, 21906769, 22278994, 35696286, 22278998, 265020, 265021, 265007, 265008, 264636, 52644150, 33657023, 264692, 264693, 29331822, 29331824, 55812038, 83373044, 56182181, 60424268, 66714117, 29331825, 33657109, 29331826, 27486262, 33657349, 56526486, 265018, 265019, 22279002, 264448, 29331830, 66712502, 264909
kinase	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED			nuclease		UNCLASSIFIED	kinase	synthase	isomerase	ļuļ
Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain							Contains protein domain (PF00989) - UNCLASSIFIED PAS domain			Contains protein domain (PF01443) - nuclease	Viral (Superfamily 1) RNA helicase					Contains protein domain (PF00641) - Inf Zn-finger in Ran binding protein and others.
79250602 (1203, 1204) Novel Protein sim. GBank gij3522961jgbjAAC34243.1 - IAC004411) putative pio kinase fArabidopsis thalianal			20436657 (1209, 1210) Novel Protein sim. GBank gij1175322 sp P44917 Y883_HAEIN - HYPOTHETICAL PROTEIN HI0883	80334582 (1211, 1212) Novel Protein sim. GBank gi 5020264 gb AAD38043.1 AF15136 - (AF151363) Cdc42 GTPase-activating protein [Mus musculus]	188664 (M74027) - mucin		80084775 (1217, 1218) Novel Protein sim. GBank gi[2496701 sp P55552 Y4LL_RHISN - HYPOTHETICAL 91,8 KD PROTEIN Y4LL			95287851 (1223, 1224) Novel Protein sim. GBank gij1877366jembjCAB07118j -	(Z92772) recD (Mycobacterium tuberculosis)		79969348 (1227, 1228) Novel Protein sim. GBank gi 5114231[gb AAD40238.1 AF13670 - (AF136709) histidine kinase YvcG (Staphylococus aureus)	39586996 (1229, 1230) Novel Protein sim. GBank gij1339950 dbjjBAA12741 - (D85230) large subunit of NADH-dependent glutamate svrittase (Plectonema borvanum)	20465331 (1231, 1232) Novel Protein sim. GBank gij544367isp P35673 GALE_ERWAM - UDP-GLUCOSE 4- EPIMERASE (GALGOVALDENASE) (UDP- GAI ACTOSE 4.EPIMERASE)	GALACTOSE 4-EPIMERASE) 91227222 (1233, 1234) Novel Protein sim. GBank 9124980971splQ60769JTNP3_MOUSE - TUMOR NECROSIS FACTOR, ALPHA-INDUCED PROTEIN 3 (PUTATIVE DNA BINDING PROTEIN A20) (ZINC FINGER PROTEIN A20)
79250602 (1203, 1204)	11466067 (1205, 1206)	81675420 (1207, 1208)	20436657 (1209, 1210)	80334582 (1211, 1212)	95361506 (1213, 1214)	11810888 (1215, 1216)	80064775 (1217, 1218)	79629413 (1219, 1220)	87586205 (1221, 1222)	95287851 (1223, 1224)		7523475 (1225, 1226)	79969348 (1227, 1228)	39586996 (1229, 1230)	20465331 (1231, 1232)	91227222 (1233, 1234)
209	†	8		909	209	Г	609	Т	911	612		813	614	615	616	617

ă	20612842 (1236 4226)	Married Control Contro			
;	20022043 (1233, 1230	(1233, 1233, 1239) Indver Protein Sim. GBank gij3439386jembjCAB5U/46.1 j (AL098839) putative aminotransferase (Streptomyces (coeficolor)		isomerase	264603
619	91227224 (1237, 1238)				56994075, 29331826, 33656970, 265008, 33657402, 33109954, 87168559, 264448, 18108374, 83373044
950	81183143 (1239, 1240,	81183143 (1239, 1240) Novel Protein sim. GBank gil464335[sp Q05922]DUS2_MOUSE - DUAL SPECIFICITY PROTEIN PHOSPHATASE 2 (DUAL SPECIFICITY PROTEIN PHOSPHATASE PAC-1)		phosphatase	29146498, 264758, 264369, 29148627
621	80239251 (1241, 1242)			UNCLASSIFIED	264556 264558 264639
622	20456427 (1243, 1244)	20456427 (1243, 1244) Novel Protein sim. GBank gil2633557 emb CAB13060 - (299110) yidF Bacillus subtilis		UNCLASSIFIED	264605
623	10131798 (1245, 1246)	10131798 (1245, 1246) Novel Protein sim. GBank gil1857710[gblAAB48482] - (U87224) contactin associated protein [Rattus norvegicus]	Contains protein domain (PF00054) - Iaminin Laminin G domain	laminin	264906
624	19534127 (1247, 1248)	19534127 (1247, 1248) Novel Protein sim. GBank gif1705703[splP52225[CCMF_PSEFL - CYTOCHROME C-TYPE BIOGENESIS PROTEIN CYCK		cytochrome	264596
625	13084619 (1249, 1250)	Novel Protein sim. GBank gi 2894252 emb CAA17114.1 - (AL021841) hypothetical protein Rv3342 [Mycobacterium tuberculosis]		UNCLASSIFIED	264688
626	88062603 (1251, 1252)	88062603 (1251, 1252) Novel Protein sim. GBank gil416592[splP32323]AGA1_YEAST - A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR		UNCLASSIFIED	29331822, 264905, 264908, 33657023, 33657109, 264558
627	80255457 (1253, 1254)	80255457 (1253, 1254) Novel Protein sim. GBank gij3098418 (AF040944) - P140 [Mus musculus]		UNCLASSIFIED	18108394, 264112, 264593, 265022, 264635
928	80077096 (1255, 1256)		Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264600
628	78851602 (1257, 1258)	78851602 (1257, 1258) Novel Protein sim. GBank gij1143204 (U34305) - ORF2; Method: conceptual translation supplied by author. [Shigella sonnel]		isomerase	264906, 264907
93	39565156 (1259, 1260)	39565156 (1259, 1260) Novel Protein sim. GBank gij3236368 (AF064748) - S3-12 [Mus musculus]		UNCLASSIFIED	264490
631	20598718 (1261, 1262)				263978
632	27843890 (1263, 1264)			UNCLASSIFIED	264906, 264600, 264605, 264769, 264689,
633	80477772 (1265, 1266)			UNCLASSIFIED	264769
25.0	70574506 (1267, 1268)				265019
36	7001000 (1209, 1270)			UNCLASSIFIED	264689
020	13310301 (1771, 1772)				264596, 264762, 264693

ı					DO 4000 00 1007 DO 4540 DO 4004
/28	82455796 (1273, 127	02433736 (1273, 1274) Novel Protein Sim. Obsank gije326739(emb CAB10933 - (298268) recN [Mycobacterium tuberculosis]		nuclease	264900, 264907, 264911, 264901, 264602, 264603, 264604, 264605, 18108351, 264762, 264766, 264687, 264769, 264689, 35695917, 264693, 264638, 264639, 264559, 18108385
638	14997457 (1275, 127	14997457 (1275, 1276) Novel Protein sim. GBank gild678662 emb CAB41074.1 - (AL049645) putative large ATP-binding protein [Streptomyces coelicolor]			264636
639	80204210 (1277, 127			struct	264112, 263974
640	17929579 (1279, 128	17929579 (1279, 1280) Novel Protein sim. GBank gi[1432083 (U60981) - homolog	Contains protein domain (PF01466) - rnapolymerase	rnapolymerase	265009, 265010
		to Skp1p, an evolutionarily conserved kinetochore protein in Skp1 family budding yeast [Arabidopsis thailana]			
<u>8</u>	79636398 (1281, 1282)	-		UNCLASSIFIED	264693
642	19898737 (1283, 1284)	84)		UNCLASSIFIED	264565
643	81516220 (1285, 1286)	(98		UNCLASSIFIED	264906, 264908, 264758, 264288, 264632, 264635, 264639, 264564
44	11751367 (1287, 1288)	88)		UNCLASSIFIED	264684
645	95010907 (1289, 1290)	[(06		UNCLASSIFIED	264906, 264762, 264693, 264639, 264559
646	80069083 (1291, 1292)	92)			264595, 264566
647	80257085 (1293, 129	_	Contains protein domain (PF00023) - transcriptfactor	transcriptfactor	264909, 264591
		Ξ	Ank repeat		
648	80077428 (1295, 129	80077428 (1295, 1296) Novel Protein sim. GBank gi 1044963 bbs 169646 - protamine [Monodonta turbinata, gonads, Peptide, 106 aa]		UNCLASSIFIED	264600
649	80247447 (1297, 1298)			UNCLASSIFIED	263978
650	11798316 (1299, 1300)	(00		UNCLASSIFIED	264686
651	11776932 (1301, 130	11776932 (1301, 1302) Novel Protein sim. GBank gij1346916 spiP12283 pURA_ECOL! - ADENYLOSUCCINATE SYNTHETASE (IMP-ASPARTATE	ш		264602, 264638
653	85516704 (1303 130	LIGASE)		UNCLASSIFIED	264905, 264907, 264909, 263978, 264637
653	82124947 (1305, 130	82124947 (1305, 1306) Novel Protein sim. GBank		UNCLASSIFIED	22278996, 264510, 264511, 264512, 264593,
}		gij17229771spjQ106381Y03C_MYCTU - HYPOTHETICAL 82.8 KD PROTEIN CY130.12C			21906754, 264603, 264760, 18108376, 264556
654	95010589 (1307, 1308)	_		UNCLASSIFIED	264906, 264595, 264632
655	79320692 (1309, 131	79320692 (1309, 1310) Novel Protein sim. GBank gij130327jspjP26647jPLSC_ECOLI - 1-ACYL-SN-GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE (1-AGPACYLTRANSFERASE) (1-AGPAT) (LYSOPHOSPHATIDICACID ACYLTRANSFERASE) (LPAAT)	Contains protein domain (PF01553) - (transferase Acyltransferase	(ransferase	264592
929	80416739 (1311, 131	12)		UNCLASSIFIED	264602, 264605, 264766, 264691
657	20611010 (1313, 1314)	14)		UNCLASSIFIED	264557, 264558

	87761815 (1315, 1316)	5689493 dbj BAA83030.1 - in [Homo sapiens]		UNCLASSIFIED	22278996, 60432049, 29331822, 29331824, 29331828, 265007, 265009, 33657402, 33657084, 265017, 264448, 21906765, 21906766, 263967, 20281149, 18108370, 18108374, 264482
629	87718663 (1317, 1318	87718663 (1317, 1318) Novel Protein sim. GBank gi[2137872 pir 148724 - zinc finger protein PZF - mouse	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	22278999, 60432049, 66714117, 29331827, 265007, 264766, 56181562, 18108359, 18108358, 4810836, 48108384
099	81897922 (1319, 1320)	0		UNCLASSIFIED	264757
681	80026023 (1321, 1322	80026023 (1321, 1322) Novel Protein sim GBank	Contains profein domain (PF00874) -	- LINCLASSIFIED	264510 265009 264600 264602 264603
		Y_BACSU - LEVANSUCRASE SIS OPERON	Transcriptional antiterminator bglG family		264604, 264605, 32833986, 18108376, 264636, 18108387, 22279000
662	20463731 (1323, 1324	20463731 (1323, 1324) Novel Protein sim. GBank gil4545229[gblAAD22450.1]AF11618 - (AF116183) SecA homolog [Actinobacillus actinomycetemcomitans]		UNCLASSIFIED	264605
663	20628080 (1325, 1326	20628080 (1325, 1326) Novel Protein sim. GBank gil5689250 dbj BAA82881.1 - (AB024335) similar to orf5 (Comamonas testosteroni)		dehydrogenase	264605
664	80508512 (1327, 1328	80508512 (1327, 1328) Novel Protein sim. GBank gi 1652848 db BAA17766 - (D90909) DNA photolyase [Synechocystis sp.]		UNCLASSIFIED	264769
999	80079053 (1329, 1330			isomerase	264600
		gij 1684 ispjr 1840icobj. PSEDE - PRECORRIN-3B CTA METHYLTRANSFERASE (PRECORRIN-3 METHYLTRANSFERASE) (PRECORRIN-3 METHYLASE)			
999	79603142 (1331, 1332	79603142 (1331, 1332) Novel Protein sim. GBank gij3261829jemb CAB10927] - (Z98260) hypothetical protein Rv1230c (Mycobacterium tuberculosis]		glycoprotein	264907, 265007
299	94631802 (1333, 1334	94631802 (1333, 1334) Novel Protein sim. GBank gil5688851ldbjlBAA82702.1 - (AB017438) Orf5 [Streptomyces coelicolor]		UNCLASSIFIED	264689, 264602, 264593
899	82051891 (1335, 1336	_	Contains protein domain (PF00453) - nbosomalprot	ribosomalprot	264905, 264906, 264908, 264600, 264601,
		(AL031541) 50S ribosomal protein L20 (Streptomyces coeticolor)	Ribosomal protein L20		264603, 264605, 264760, 264689, 264636. 264638, 264639
699	12967154 (1337, 1338)			UNCLASSIFIED	264637
670	80238549 (1339, 1340	80238549 (1339, 1340) Novel Protein sim. GBank gi 2582531 (AF026444) - 2- Isopropylmalate synthase [Streptomyces coelicolor]		synthase	264905, 264906, 264908, 264601, 264762, 264766, 264689, 264638, 18108385, 264486
671	79601368 (1341, 1342)		Contains protein domain (PF00023) - UNCLASSIFIED Ank repeat	UNCLASSIFIED	264690, 264692, 264693, 264636, 18108387
672	79834371 (1343, 1344	79834371 (1343, 1344) Novel Protein sim. GBank gil2114430 (U92703) - Olf-1/EBF- like-3 transcription factor [Mus musculus]		transcriptfactor	264910, 265017
673	82285798 (1345, 1346	82285798 (1345, 1346) Novel Protein sim. GBank gil4589285 gb AAD26430.1 AF13515 - (AF135154) ferric alcaligin siderophore receptor [Bordetella pertussis]			264759
674	79199259 (1347, 1348)			UNCLASSIFIED	264629

				ſ	
6/2	8/8958/0 (1349, 1350)	8789367U (1349, 1350) Novei Protein Sim. Gbank gil4980755 gb AAD35347.1 AE00170 - (AE001708) D-	Contains protein domain (Pro1620) - UNCLASSIFIED D-ala D-ala ligase		204400, 22270999, 00714117, 204300, 264511, 265008, 60433438, 264600, 264601.
		alanineD-alanine ligase [Thermotoga maritima]	,		284602, 264603, 264604, 264605, 264762.
					264687, 264769, 60431602, 18108374,
978	70900007 (1361 1360)	70900607 (1361 1360) Namel Bestein cim CBent			266010
0	1,0033007 (1,331, 1,332)	Novel Protein Sim. Gbank pil1723566sp0010479IYDF7 SCHPO - PUTATIVE			0.000
		GLUCOSYLTRANSFERASE C17C9.07			
229	21644312 (1353, 1354)	21644312 (1353, 1354) Novel Protein sim. GBank gil687208 (U03976) - dynein		ATPase_associated	264591, 264632
		neavy chain isotype oc i i npheustes granual			F. 20
878	84225200 (1355, 1356)	84225200 (1355, 1356) Novel Protein sim. GBank gi 1586274 pri 2203365A - aminicalabas Maria	Contains protein domain (PF00053) - Iaminin	laminin	264758, 264682, 264557
679	79868855 (1357, 1358)	79868855 (1357, 1358) Novel Protein sim. GBank gi[3928723 emb CAA22219 -		UNCLASSIFIED	22278996, 264693
		(AL034355) putative ABC transporter (Streptomyces			
		coelicolor]			
989	20726424 (1359, 1360)				264600, 264602
189	94322017 (1361, 1362)	94322017 (1361, 1362) Novel Protein sim. GBank	Contains protein domain (PF00053) - Iaminin	laminin	264102, 264907, 264908, 265006, 264693,
		gij5174493 ref NP_006050.1 pLAMC - Iaminin, gamma 3	Laminin EGF-like (Domains III and		263972, 83373044, 264566
			S		
682	11392476 (1363, 1364)			ED	264595
683	80083680 (1365, 1366)	80083680 (1365, 1366) Novel Protein sim. GBank	Contains protein domain (PF00782) - phosphatase		264634
		gil4758208 ref NP_004081.1 pDUSP - dual specificity	Dual specificity phosphatase,		
		phosphatase 3 (vaccinia virus phosphatase VH1-related)	catalytic domain		
684	20465367 (1367, 1368)	20465367 (1387, 1368) Novel Protein sim. GBank gil5420387lemb CAB46679.1 -			264605
		(AJ243459) proteophosphoglycan (Leishmania major)			
685	80246735 (1369, 1370)				264909, 263967, 263981
989	79208606 (1371, 1372)				264631
687	80085629 (1373, 1374)				264693, 264635
689	79853412 (1375, 1376)	79853412 (1375, 1376) Novel Protein sim. GBank gi 2688962 (AF027768) - LspA		peptidase	264907, 264638
		(Serratia marcescens)			
689	88064256 (1377, 1378)	88064256 (1377, 1378) Novel Protein sim. GBank gij3046931 (AF049330) - PPAR	Contains protein domain (PF00076) - UNCLASSIFIED	UNCLASSIFIED	264906, 264907, 265007, 265009, 60433438,
		gamma coactivator (Mus musculus)	RNA recognition motif. (a.k.a. RRM,		21906754, 264760, 18108358, 21906766,
			RBD, or RNP domain)		21906769, 265021, 18108361, 263974,
					18108379, 264557, 18108385, 22279002
069	80389750 (1379, 1380)	80389750 (1379, 1380) Novel Protein sim. GBank		UNCLASSIFIED	264510, 264511, 264764, 264769
		gij2498941 sp Q15428 SP62_HUMAN - SPLICEOSOME associated Protein 82 (sab 62) (se3a66)			
991	81854392 (1381 1382)			UNCLASSIFIED	264757
692	83608936 (1383, 1384)	83608936 (1383, 1384) Novel Protein sim. GBank qil5420387 emblCAB46679.11 -	Contains protein domain (PF00097) - UNCLASSIFIED		55812038, 55811957, 265018, 55811150,
_	•		Zinc finger, C3HC4 type (RING finger)		18108351, 264908, 60431528, 264594
693	79586116 (1385, 1386)	79586116 (1385, 1386) Novel Protein sim. GBank oil854065lemblCAA583371 -		UNCLASSIFIED	264635
		(X83413) U88 [Human herpesvirus 6]			
694	82455983 (1387, 1388)	82455983 (1387, 1388) Novel Protein sim. GBank			22278996, 264510, 264602, 264603, 264762.
		gi 267327 sp Q01033 VG48_HSVSA - HYPOTHETICAL GENE 48 PROTEIN			264687, 264769, 264693

	3, 31, 11, 5			,, 0,0000000000000000000000000000000000	Chichook Iolan I woodchor	01111004 10141	TO0100 30100 00100 3E300103
2					von Willebrand factor type D domain		29331830, 264908, 264909, 264511, 265007, 264910, 264758, 264764, 264288, 65274791
969	79830982 (1	1391, 139	<u>.</u> 9		Contains protein domain (PF00005) - transport ABC transporter	transport	264905, 264595
269	11767889 (1	1393, 139	11767889 (1393, 1394) Novel Protein sim. GBank gi 1731343 sp Q10694 yY25_MYCTU - HYPOTHETICAL 24,4 KD PROTEIN CY49.25		Contains protein domain (PF01836) - UNCLASSIFIED Transposase	UNCLASSIFIED	264682
869	66695862 (1395, 1396)	1395, 139	(9)			UNCLASSIFIED	264688, 35695917
669	79582558 (1397, 1398)	1397, 139	(8)			UNCLASSIFIED	264682
700	79639098 (1399, 1400)	1399, 140					264693
701	80230242 (1	1401, 140	80230242 (1401, 1402) Novel Protein sim. GBank gi 100 {(D64003) hypothetical protein {S	1001236 dbj BAA10477 - Synechocystis sp.]	_	UNCLASSIFIED	264488, 264510, 264511, 264602, 264605, 264689
702	79814789 (1403, 1404)	1403, 140	l			UNCLASSIFIED	264909
703	20446820 (1	1405, 140	20446820 (1405, 1406) Novel Protein sim. GBank gi[2498935 sp Q46338 SOXG_CORSP - SARCOSINE OXIDASE GAMMA SUBUNIT			oxidase	264604
\$	94312224 (1	1407, 140	94312224 (1407, 1408) Novel Protein sim. GBank gil3150513 (AF067219) - contains similarity to the kelch/MIPP family [Caenorhabditis elegans]		Contains protein domain (PF01344) - UNCLASSIFIED Kelch motif	UNCLASSIFIED	264288, 56181562, 33657109, 264629. 55811576
705	17932141 (1	1409, 141	otein sim. GBank gil ical protein o206 - E	421091 pir S30730 - scherichia coli		UNCLASSIFIED	265006
902	20288062 (1	1411, 141	20288062 (1411, 1412) Novel Protein sim. GBank gil3024872lsplQ55790lY074 SY	SYNY3 - HYPOTHETICAL			264600
			1				
707	20638065 (1	1413, 141	20638065 (1413, 1414) Novel Protein sim. GBank gil342 (AF075709) ABC transporter AT [Pseudomonas putida]	3420608 gb AC31907.1 - ATP-binding subunit		transport	264603
8 9	20708292 (1415, 1416)	1415, 141	_				264601, 264692
709	88001439 (1417, 141	88001439 (1417, 1418) Novel Protein sim. GBank gij36497 [(AJ000281) mucin [Homo sapiens]	3649741[emb CAA03985 - piens]		struct	18108398, 264637, 264908, 264909
710	11356683 (1419, 142	11356683 (1419, 1420) Novel Protein sim. GBank gi 3080425 emb CAA18744.1 - [AL022604) putative protein [Arabidopsis thaliana]	3080425[emb[CAA18744.1] - Arabidopsis thaliana]			264369
711	17931418 (1421, 1422)	1421, 142	2)				265019
712	80258164 (1423, 144	80258164 (1423, 1424) Novel Protein sim. GBank gil4758686jrefiNP_002323.1pLRP1 - low density related protein 1 (alpha-2-macroglobulin receptor)	lipoprotein	Contains protein domain (PF00058) - apolipoprotein pLRP1 - fow density lipoprotein Low-density lipoprotein receptor repeat class B	apolipoprotein	284591
713	79263126 (1425, 142	79263126 (1425, 1426) Novel Protein sim. GBank gil1703266 splQ11056 AMI2_MYCTU - PUTATIVE AMIDASE CY50.19C	YCTU - PUTATIVE		hydrolase	264906, 264907
714	27847651 (1427, 14	Bank 11692.1 ransfer	pBAAT - bile acid Coenzyme ase; glycine N-			264508, 264555

1	70030470 4420	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		0010100410141	200700	_
2	1.9039423 (1428, 1430)	hypothetical protein [Escherichia coli]		UNICHONIC	708407	
716	79559072 (1431, 1432)				264692	_
717	79491842 (1433, 1434)	79491842 (1433, 1434) Novel Protein sim. GBank		dehydrogenase	264636	_
		gi[2494074[sp[P55653]GABD_RHISN - PROBABLE SUCCINATE-SEMIALDEHYDE DEHYDROGENASE				
		(NADP+) (SSDH)				_,
718	94319658 (1435, 1436)		Contains protein domain (PF00093) - kinase	kinase	18108392, 22278994, 22278998, 265008,	
		(Z71178) similar to pro-collagen domains; cDNA EST	von Willebrand factor type C domain		265018, 264681, 18108354, 264684, 264685,	
		EMBL: U2/9/8 comes from this gene; cDNA ES			264686, 264687, 264689, 21906769,	
		EMBL: D27977 comes from this gene; cDNA EST			18108361, 264691, 264692, 55810764,	
_		EMBL:D34199 comes from this gene; cDNA EST			264635, 18108381, 18108382, 83373044,	_
		EMBL:D64392 comes from this gene; cDNA EST EMBL			18108388	٠-,
719	17679564 (1437, 1438)	Novel Protein sim. GBank gi[2104302 emb CAB08631] -		UNCLASSIFIED	265011	
		(295387) hypothetical protein Rv2611c [Mycobacterium				
		[tuberculosis]				_
720	79841684 (1439, 1440)				264908	-
721	15020180 (1441, 1442)	15020180 (1441, 1442) Novel Protein sim. GBank		UNCLASSIFIED	264629	
		gij123530 sp P04929 HRPX_PLALO - HISTIDINE-RICH				
		GLYCOPROTEIN PRECURSOR				
722	9862603 (1443, 1444)	Novel Protein sim. GBank gil498253 (U02372) - integrase			264910	
		[Vibrio cholerae]				
723	19755599 (1445, 1446)	Novel Protein sim. GBank gi[2253054 emb CAB10705 -		UNCLASSIFIED	264691	_
724	10126494 (1447, 1448)	10126494 (1447, 1448) Novel Protein sim. GBank gil4063015 (AF083061) -	Contains protein domain (PF00353) - protease	protease	264909	_
			Hemolysin-type calcium-binding proteins			
725	78878679 (1449 1450)			UNCLASSIFIED	264905, 264907	_
728	13086282 (1451, 1452)			1	264636	_
727	13522872 (1453, 1454)			l	264634	_
728	20268471 (1455, 1456)	20268471 (1455, 1456) Novel Protein sim. GBank gil2633910 emb CAB13411 -			264567	
		(Z99112) similar to hypoth				_
729	11293753 (1457, 1458)			UNCLASSIFIED	264490	_
730	19900373 (1459, 1460)			isomerase	264564	_
		gij2494660 sp Q45291 GALE_BRELA - UDP-GLUCOSE 4-				
		EPIMERASE (GALACTOWALDENASE) (UDP.				_
		GALACTOSE 4-EPIMERASE)				_
731	80058750 (1461, 1462)	80058750 (1461, 1462) Novel Protein sim. GBank gil1146192 (L47838) - putative		UNCLASSIFIED	264605	
		[Bacillus subtilis]				T
732	80258175 (1463, 1464)	80258175 (1463, 1464) Novel Protein sim. GBank		struct	264591, 264594, 264595	_
		[gi[1168396 sp P46681 AIP2_YEAST - ACTIN				_
						-
733	20446839 (1465, 1466)			UNCLASSIFIED	264604	1
734	20435987 (1467, 1468)	20435987 (1467, 1468) Novel Protein sim. GBank gij3184080jembjCAA19336j -		ubiquitin	264604	
		(AL023781) hypothetical protein [Schizosaccharomyces				
		[bombe]				7

					264504
735	11607959 (1469, 1470)	11607959			
736	10879734 (1471, 1472)	10879734 (1471, 1472) Novel Protein sim. GBank gil400831[sp P31135 POTH_ECOLI - PUTRESCINE TRANSPORT SYSTEM PERMEASE PROTEIN POTH	Contains protein domain (PF00528) - Is Binding-protein-dependent transport systems inner membrane component	transport	264636
737	78945340 (1473, 1474)		Contains protein domain (PF00815) - UNCLASSIFIED Regulator of G protein signaling domain		265020
738	17895353 (1475, 1476)				265008
739	79833670 (1477, 1478)	79833670 (1477, 1478) Novel Protein sim. GBank gi[2506867[splp33225]TORA_ECOLI - TRIMETHYLAMINE- N-OXIDE REDUCTASE PRECURSOR (TMAO REDUCTASE) (TRIMETHYLAMINE OXIDASE)	<u> </u>	oxidase	264910
240	10881557 (1470 1480)				264907, 264764, 264634, 264637
1	79827273 (1481, 1482)	79827273 (1481, 1482) Novel Protein sim. GBank gij3261828 emb CAB10925 - 1298260) mm [Mycobacterium tuberculosis]	Contains protein domain (PF01883) - UNCLASSIFIED Domain of unknown function	UNCLASSIFIED	264689, 35696286, 264510, 264908, 18108362
742	82393795 (1483, 1484)	82393795 (1483, 1484) Novel Protein sim. GBank gij3877494 emb CAA88472.1 - (248563) ATP binding protein with similarity to the CDC48/PAS1/SEC18 family; cDNA EST EMBL:D65037 comes from this gene; cDNA EST EMBL:D68340 comes from this gene; cDNA EST EMBL:D65048 comes from this gene; cDNA EST EMBL:D65048 comes from this		UNCLASSIFIED	29331822, 264910, 264762
		gene; cund Es i EMBL: U0043	Contains acatain domain (PE00145)	UNC! ASSIFIED	264488 264259, 264508, 264905, 264906,
	82300051 (1485, 1486)	82300051 (1485, 1486) Novel Protein sim. GBank gi 127420jsp P19888 MTBA_BACAR - MODIFICATION METHYLASE BANI (CYTOSINE-SPECIFIC METHYLTRANSFERASE BANI) (M.BANI)	C-5 cytosine-specific DNA methylase		26407, 264906, 264909, 264510, 264511, 264512, 265008, 265009, 264910, 264591, 264566, 264759, 265010, 265011, 18108351, 264763, 264766, 264766, 264768, 264630, 18108370, 264629, 18108372, 264630, 264631, 264634, 264558, 18108385, 264482, 264564, 264567
744	80230421 (1487, 1488)				18108397, 264511, 264690, 264628, 264638, 264639, 264692, 264639, 264766
745	9841963 (1489, 1490)	Novel Protein sim. GBank gi 78921 pir S04846 - UDP-N-acetylmuramoylalanyl-D-glutamyl-2, 6-diaminopimelateD-alanyl-D-alanine ligase (EC 6.3.2.15) precursor - Fscherichia coli		glycoprotein	264906
746	11073229 (1491, 1492)	11073229 (1491, 1492) Novel Protein sim. GBank gil3386354 (AF074705) - nvorchelin svnihelase iPseudomonas aerudinosal		synthase	264600
747	94322044 (1493, 1494)	94322044 (1493, 1494) Novel Protein sim. GBank gi 2887411 db BAA24848 - (AB007878) KIAA0418 [Homo sapiens]	Contains protein domain (PF00018) - oxidase SH3 domain	oxidase	66714117, 264905, 264509, 264906, 264907, 264908, 264908, 264911, 264910, 265011, 264681, 264288, 264766, 264687, 264769, 21906768, 35695917, 264691, 264693, 264634, 264634, 264635, 264639, 26182323, 83373044
748	11617923 (1495, 1496)				264690

- 1	20469119 (1497, 1498)			UNCLASSIFIED	264604
6707	0427 (1499, 1500)	COCSOCALI (1499, 1500) Novel Protein sim. GBank gild 169727 jsp P44948 FPG_HAEIN - FORMAMIDOPYRIMIDINE-DNA GLYCOSYLASE (FAPY-			264600
á S	36169 (1501, 1502)	Novel Protein sim. GBank gl[5360068]gbJAAD42851.1[AF15968 - (AF159689) serine/threonine kinase PKN3 [Myxococcus xanthus]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264605, 264559
824	50366 (1503, 1504)	82450366 (1503, 1504) Novel Protein sim. GBank gil1168662[sp P44426 BIOA_HAEIN - ADENOSYLMETHIONINE-8-AMINO-7-OXONONANOATE AMINOTRANSFERASE (7,8-DIAMINO-PELARGONIC ACID AMINOTRANSFERASE) (DAPA AMINOTRANSFERASE)	Contains protein domain (PF00202) - Aminotransferases class-III pyridoxal phosphate	UNCLASSIFIED	264508, 264907, 264510, 265011, 264762, 264689, 35695855, 264638, 18108387
805		80508718 (1505, 1506) Novel Protein sim. GBank gi[2851530]sp P32399 YHGE_BACSU - HYPOTHETICAL 84.1 KD PROTEIN IN HEMY-GLTT INTERGENIC REGION (ORFB)		UNCLASSIFIED	264909, 264600, 264602, 264604, 264760, 264769, 264634
950	95083741 (1507, 1508)			UNCLASSIFIED	264508, 264906, 264907, 264908, 264909, 264759, 264602, 264764, 264769, 264628, 264629, 264629, 264630, 264632, 264634, 264639, 264638, 264648, 264688
8	35449 (1509, 1510)	80185449 (1509, 1510)		UNCLASSIFIED	264448, 264690
<u>ģ</u>	31086 (1511, 1512)	Novel Protein sim. GBank gij3449276jemb CAA20420j - (AL031317) putative dehydrogenase [Streptomyces coelicolor]			264769, 264689, 264638, 264639
	19400333 (1513, 1514)			UNCLASSIFIED	264682 264685
	33176 (1515, 1516)	/8963176 (1515, 1516) Novel Protein sim. GBank gil4580331 emb CAB40107.1 - (AJ001206) putative glycogen debranching enzyme (Streptomyces coelicolor)		amylase	265007, 18108387, 265007, 18108387
\$ 18	(1517, 1518)	[2911858 (AF047659) - No habditis elegans]		UNCLASSIFIED	264684, 264686
	10201, 1520)	o/oz8888 (1519, 1520) Novel Protein sim. GBank gij3451312 emb CAA20449j - (AL031324) membrane atpase (Schizosaccharomyces pombej	Contains protein domain (PF00122) - transport E1-E2 ATPase	ransport	29331822, 264908, 52644045, 56182435, 60170831, 21906754, 265017, 265019, 264681, 264687, 264688, 21906766, 21906768, 21906768, 265020, 265021, 265022, 264635,
8	79877966 (1521, 1522)			UNCLASSIFIED	2527 9000 264766
8002	(3563 (1523, 1524)	80023563 (1523, 1524) Novel Protein sim. GBank gil3327158(dbj BAA31647 - (AB014572) KIAA0572 protein [Homo sapiens]			264907, 264593, 265020
		Novel Protein sim. GBank gl 4981266 gb AAD35922.1 AE00174 - (AE001744) lipopolysaccharide core biosynthesis protein KdtB [Thermotoga maritima]			264600
383	39515024 (1527, 1528)				264603

765	80025347 (1529, 1530)	80025347 (1529, 1530) Novel Protein sim GBank nil3845093 (AF001371) .		to to	26400E 26400E 264E04 264686 2262202
		erythrocyte membrane protein PfEMP3 [Plasmodium		15016	204303, 204300, 204334, 204000, 33037023
		falciparum]			
992	82417404 (1531, 1532)				264605, 264762, 18108374
787	10296742 (1533, 1534)	10296742 (1533, 1534) Novel Protein sim. GBank gij541121 pir S40827 - hypothetical protein 0300 - Escherichia coli		UNCLASSIFIED	264907
89/	79416080 (1535, 1536)			UNCLASSIFIED	264592, 264595
169	80086554 (1537, 1538)	80086554 (1537, 1538) Novel Protein sim. GBank gi[2982501 emb CAA06164 - (AJ004832) neuropathy target esterase [Homo sapiens]		esterase	55810764, 264559
770	80417847 (1539, 1540)	80417847 (1539, 1540) Novel Protein sim. GBank gij283437 pir S27850 -		UNCLASSIFIED	264905, 264907, 264828, 264909, 265010
		hypothetical protein - Trypanosoma cruzi (fragment)			264766, 264628, 264629, 264634, 264636, 264555
771	95329509 (1541, 1542)		Contains protein domain (PF00097) - UNCLASSIFIED	UNCLASSIFIED	56182575, 35696286, 264259, 66714117,
		Ξ	AF14059 - (AF140598) ring-box Zinc finger, C3HC4 type (RING		264107, 66712502, 56182435, 264112,
		protein 1 [Homo sapiens]	finger)		55812038, 87168559, 264288, 21906766, 33657023, 65274620, 65274791, 18108381
772	78971362 (1543, 1544)			UNCLASSIFIED	264910
773	78945363 (1545, 1546)			UNCLASSIFIED	265020
424	79856129 (1547, 1548)	79856129 (1547, 1548) Novel Protein sim. GBank gil5531324 emblCAB51045.1 -		UNCLASSIFIED	264909
		(AJUUS)/9) purative aikane 1-monooxygenase [Pseudomonas fluorescens]			
175	20620141 (1549, 1550)	-		UNCLASSIFIED	264555
9//	78942693 (1551, 1552)		Contains protein domain (PF01006) -		265019
			Hepatitis C virus non-structural protein NS4a		
111	79960378 (1553, 1554)	79960378 (1553, 1554) Novel Protein sim. GBank	Contains protein domain (PF01344) - protease	protease	21906754, 265020, 60170615, 264691
		gil4505461 ref NP_003624.1 pNRPB - nuclear restricted protein, BTB domain-like (brain)	Kelch motif		
778	20691310 (1555, 1556)			UNCLASSIFIED	264511
779	80054024 (1557, 1558)				264603
780	95288987 (1559, 1560)	95288987 (1559, 1560) Novel Protein sim. GBank gil1144520 (U34956) -		synthase	264907, 264600, 264601, 264602, 264603,
		phosphoribosyfformyfglycinamidine synthase [Mycobacterium tuberculosis]			264604, 264605, 264486
781	80250049 (1561, 1562)				264905, 264907, 265010, 264600, 264601, 18108362, 18108374, 264556
782	8758529 (1563, 1564)	Novel Protein sim. GBank gil4155447 (AE001517) - proline/hetaine transporter (Helicoharter polori 1991		UNCLASSIFIED	264605
783	16410791 (1565, 1566)				265020
784	80051197 (1567, 1568)			UNCLASSIFIED	264635, 33657023, 29331828, 265017,
į					264565, 264566
785	56073541 (1569, 1570)	56073541 (1569, 1570) Novel Protein sim. GBank gi[3451335 (AC005525) - F22162_1 [Homo sapiens]	Contains protein domain (PF00047) - struct Immunoglobulin domain	struct	35696052, 264604
98/	20438842 (1571, 1572)			transport	264603
		giji36748ispip10905juGPA_ECOLI - SN·GLYCEROL·3- PHOSPHATE TRANSPORT SYSTEM PERMEASE PROTEIN UGPA			
787	80258364 (1573, 1574)			UNCLASSIFIED	264593

788	80507844 (1575, 157	80507844 (1575, 1576) Novel Protein sim. GBank gil2746079 (AF015310) - BTH1		synthase	264909, 264602, 264603, 264769, 264638
789	17294715 (1577, 157	17294715 (1577, 1578) Novel Protein sim. GBank gil2351849 (U93357) - 40 kDa heat shock chaperone ordein [Halobacterium culimbrum]		UNCLASSIFIED	265007
790	86284406 (1579, 156	86284406 (1579, 1580) Novel Protein sim. GBank gil5706378 dbjjBAA83099.1 - (AB026118) MALT1 [Homo sapiens]	Contains protein domain (PF00047) - glycoprotein Immunoglobulin domain	glycoprotein	22278999, 29331824, 264828, 87188559, 265018, 21906765, 21906767, 21906768, 21906769, 264692, 22279000, 264563
791	94651627 (1581, 156	94651627 (1581, 1582) Novel Protein sim. GBank gil5689948lemb CAB51985.1 - (AL109663) putative isoleucyl-tRNA synthetase (Streptomyces coelicolor A3(2))			264601, 264605, 284636
792	80058786 (1583, 158	80058786 (1583, 1584) Novel Profein sim. GBank gi[393194 (L02375) - S-antigen [Plasmodium falciparum]		struct	265021, 264631, 264635, 264556
793	79638730 (1585, 158	79638730 (1585, 1586) Novel Protein sim. GBank gi 1345408 dbj BAA05046 - (D26046) AT motif-binding factor [Mus musculus]	Contains protein domain (PF00046) - homeobox Homeobox	homeobox	264693
794	81839294 (1587, 158	81839284 (1587, 1588) Novel Protein sim. GBank gi†105884 pir S24023 - dopamine receptor D4 - human (fragment)		UNCLASSIFIED	264603, 264604, 264910, 264762, 264906, 264639, 264909, 264757
795	80074988 (1589, 156	80074988 (1589, 1590) Novel Protein sim. GBank gil 1877334 emb CAB07082 - (292771) birA [Mycobacterium tuberculosis]		carboxylase	264488, 35696052, 264905, 264907, 265010, 35696423, 264636
796	86669451 (1591, 1592)				60432229, 55811150, 264630, 264637, 264565
797	87771781 (1593, 156	87771781 (1593, 1594) Novel Protein sim. GBank gil2995447 emb CAA71519 - (Y10495) CDV-1R protein [Mus musculus]		struct	22278998, 264093, 264094, 66714117, 21906767, 21906769, 265020, 265022
798	79865209 (1595, 1596)			transcriptfactor	264687, 264768, 264693
789	79557816 (1597, 159	79557816 (1597, 1598) Novel Protein sim. GBank gil4467250 emb CAB37575 - (AL035569) probable Glu-IRNA Gln amidotransferase subunit [Streptomyces coelicolor]		hydrolase	264909, 264910, 264636, 264638
800	79970189 (1599, 1600)			UNCLASSIFIED	264488
108	80499399 (1601, 16	80499399 (1601, 1602) Novel Protein sim. GBank gil2791517 emb CAA16054 - (AL021246) hypothetical protein Rv2477c [Mycobacterium tuberculosis]		transport	264508, 264511, 265006, 265009, 264769, 264567, 264486
802	79834598 (1603, 160	79834598 (1603, 1604) Novel Protein sim. GBank gil4887211[gb]AAD32237.1[AF14744 - (AF147449) penicilin binding protein 1B [Pseudomonas aeruginosa]			264905, 264693
803	20467520 (1605, 1606)	-		struct	264605
80 8	10174239 (1607, 160	10174239 (1607, 1608) Novel Prolein sim. GBank gij1176152 sp P44507 YHAD_HAEIN - HYPOTHETICAL PROTEIN HI0091		kinase	264510
805	7959993 (1609, 1610)	_			264508
808	80484113 (1611, 16	80484113 (1611, 1612) Novel Protein sim. GBank gil2764612 emb CAA04683 - (AJ001330) omithine transcarbamoylase [Lactobacillus sakei]	Contains protein domain (PF00185) - transferase Aspartate/ornithine carbamoyltransferase	transferase	264769
807	80381812 (1613, 16	80381812 (1613, 1614) Novel Protein sim. GBank gij2833311[sp Q21828 YNFD_CAEEL - HYPOTHETICAL 18.9 KD PROTEIN R07E5.13 IN CHROMOSOME III			264764

909	25405047 14545 46461	4			
g	33100017 (1013, 1016)	SSTUGGET (1013, 1018) NOVER PTOTEIN GBANK gij3913092 sp Q46170 ARCD_CLOPE - ARGININE/ORNITHINE ANTIPORTER		transport	264909, 264602, 21906764, 18108374
608	81454254 (1617, 1618)	81454254 (1617, 1618) Novel Protein sim GRank	Contains and demain (BECOCTA)	001000	010100 200100 000100 000100
}		ail3913016isplP74309iALF1 SYNY3 - FRUCTOSF-	Frictose-bisphosphate aldolase	UNCLASSIFIED	264306, 264906, 264909, 265007, 264910, 1
		۱ <u>%</u>	class-I		264687, 264769, 264689, 264636, 264486
810	80192761 (1619, 1620)	80192761 (1619, 1620) Novel Protein sim. GBank	Contains protein domain (PF00248) - reductase	reductase	264369
		gij401472ispp30863jYAFB_ECOLI - HYPOTHETICAL OXIDOREDUCTASE IN ASPU-MLTD INTERGENIC REGION	Aldo/keto reductase family		
811	80079280 (1621, 1622)			UNCLASSIFIED	264558
812	10297654 (1623, 1624)			UNCLASSIFIED	264692
813	79612280 (1625, 1626)			INCI ACCIETED	264006
814	80473427 (1627, 1628)	80473427 (1627, 1628) Novel Protein sim. GBank gi[146168 (J01617) - glutaminyl- IRNA symbologia Especialis		synthase	264905, 264602, 264605, 264682, 264687,
446	05440542 (4620 4620)				204/09, 204030
2	(1601, 1631, 1631, 1631)	934 1931 3 (10.29, 10.30) Nover Protein sm. GBank gild 5995 5 2 (dB) [BAA 76848.1 - (AB02 3221) KIAA 1004 protein [Homo sapiens]		UNCLASSIFIED	264488, 22278998, 22278999, 29331822, 29331824, 29331827, 29331827, 29331828, 29331827, 29331828, 29331827, 29331828, 29346499, 264908, 265007, 265017, 265017, 265018, 265019, 264369, 264288, 264685, 21906765, 21906767, 265020, 265021, 264692, 65274620, 33657109, 264629, 18108376, 264635, 264655, 2646
818	19881910 (1631, 1632)				264600
P47	06203346 /1633 1634)	06202346 (1623 4634) Novel Dratein ein Charl Historia alle Anna Lie Anna Lie Anna Charle		11000	20000
<u>.</u>	95293510 (1055, 1054)			UNCLASSIFIED	264595
818	90938190 (1635, 1636)	90938190 (1635, 1636) Novel Protein sim. GBank gi[1477468 (U35244) - vacuolar			65274572, 22278999, 60424269, 35696052,
					55812038, 21906768, 55811957, 35695917, 33657023, 18108370, 18108374, 55810764, 35696423, 55811576, 264636
818	80254977 (1637, 1638)	80254977 (1637, 1638) Novel Protein sim. GBank gi 1001352 dbj BAA10839 - (D64006) ABC transporter [Synechocystis sp.]		transport	264565
820	80059688 (1639, 1640)	80059688 (1639, 1640) Novel Protein sim. GBank gij586814 spjP37484 YYBT_BACSU - HYPOTHETICAL		UNCLASSIFIED	264500, 264602, 264604
821	79762590 (1641, 1642)	A STAND THE PROPERTY OF THE PROPERTY PEGICINAL		INCI ASSIEIED	264910
223	80215240 (1642 1644)			OLIVOR OF THE	01000
778	60213310 (1643, 1644)			UNCLASSIFIED	264510, 264594, 264637
823	94992299 (1645, 1646)	94992299 (1645, 1646) Novel Protein sim. GBank gij3878400 emb CAA95828 -		struct	264509, 264687, 264691
		Mouse T-complex-associated-testes-expressed-1 protein			
		(PIR Acc. No. A45841); cDNA EST EMBL: D32742 comes			
		from this gene; cDNA EST EMBL: D33617 comes from this gene; cDNA EST			
824	80411171 (1647, 1648)	Novel Protein sim. GBank gi 1370076 emb CAA66887 - (X98235) type I [Drosophila melanogaster]	Contains protein domain (PF01429) - Methyl-CpG binding domain		264910, 264763, 264769, 264693

264592	264605	264603	52645080, 264508, 264905, 264509, 264906, 264907, 264907, 264908, 264909, 264512, 264910, 264591, 264591, 264591, 264591, 264600, 264766, 264768, 21906768, 35695917, 264691, 264628, 264630, 264632, 264634, 264635, 264636, 264636, 264636, 264636, 264636, 264636, 264686, 264686	264758, 264634	264603	66714117, 264910, 264639	264906	264602	264600, 264602, 264769, 264689, 264636	264768, 263994, 21906767, 264910, 264632, 264635, 264639, 264693, 83373044, 264758, 35696052, 22279002, 264508, 264905, 264906, 264448, 263972, 264908, 264909	35695917, 264557	264595	264687	264906	264762, 264556
UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	kinase		transport	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	interleukinrecept
									Contains protein domain (PF00224) - kinase Pyruvate kinase	·					Contains protein domain (PF00097) - interleukinrecept Zinc finger, C3HC4 type (RING finger)
20638600 (1649, 1650) Novel Protein sim. GBank gij3025132 sp P77391 YEAG_ECOLI - HYPOTHETICAL 74.5 KD PROTEIN IN GAPA-RND INTERGENIC REGION	11075047 (1651, 1652) Novel Protein sim. GBank gi 3242281 emb CAA16669 - (AL021646) hypothetical protein Rv3202c [Mycobacterium [tuberculosis]	80054207 (1653, 1654) Novel Protein sim. GBank gij3417424 emb CAA20312 - AL031261) putative transport protein Schizosaccharomyces pombe]	95106322 (1655, 1656) Novel Protein sim. GBank gil4336692 gb AAD17897 - (AF101361) Abnormal X segregation [Drosophila melanogaster]		20396091 (1659, 1660) Novel Protein sim. GBank gij3820539 (AF080002) - UDP-N-acetylmuramyl tripeptide synthetase MurC [Heliobacillus mobilis]		19536322 (1663, 1664) Novel Protein sim. GBank gi 1870004 emb CAB06855 - (292539) hypothetical protein Rv1024 Mycobacterium [tuberculosis]	20726654 (1665, 1666) Novel Protein sim. GBank gil2500056 sp Q46267 PFLA_CLOPA · PYRUVATE FORMATE-LYASE ACTIVATING ENZYME	21428762 (1667, 1668) Novel Protein sim. GBank gi 2497531 sp Q46078 KPYK_CORGL - PYRUVATE KINASE (PK)		66126552 (1671, 1672) Novel Protein sim. GBank gil699315 (U15184) - phosphate transport protein PSTC (Mycobacterium leprae)		79184203 (1675, 1676) Novel Protein sim. GBank gil728867 sp P40602 APG_ARATH - ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR		80059851 (1679, 1680) Novel Protein sim. GBank gil4557753 ref NP_000372.1 pMID1 - midline 1 protein
20638600 (1649, 1650)	11075047 (1651, 1652)	80054207 (1653, 1654)	9510 6322 (1655, 1656)	81742215 (1657, 1658)		[87112435 (1661, 1662)	19536322 (1663, 1664)	20726654 (1665, 1666)	21428762 (1667, 1668)	94140482 (1669, 1670)	66126552 (1671, 1672)	79450450 (1673, 1674)	79184203 (1675, 1676)	79641125 (1677, 1678)	80059851 (1679, 1680)
1	826	827	828	829	830	831	832	833	834	835	836	837	838	839	840

80078724 (1683, 1684) Novel Protein sim. GBank gil2114321 db BAA20037 - (D88733) membrane glycoprotein [Equine herpesvirus 1] 87002847 (1685, 1686) Novel Protein sim. GBank gil282325 db BAA34522.1 - (AB018345) KIAA0802 protein Homo sapiens 17941439 (1687, 1689) Novel Protein sim. GBank gil2224721 db BAA20844 - (AB002398) KIAA0390 Homo sapiens 18346844 (1689, 1690) AB002399 KIAA0390 Homo sapiens 1863341 (1681, 1682) Novel Protein sim. GBank gil2224721 db BA20844 Equialtory protein vs.m. GBank gil226579 pr A36929 - virulence regulatory protein vs.m. GBank gil226579 pr A36929 - virulence regulatory protein vs.m. GBank gil226579 pr A36929 - virulence regulatory protein sim. GBank gil224721 db A36929 - virulence regulatory protein sim. GBank gil2349 (1689, 1700) Novel Protein sim. GBank gil3499 (1689, 1700) Novel Protein sim. GBank gil3499 (1689, 1701) Novel Protein sim. GBank gil34949 (1689, 1701) Novel Protein sim. GBank gil3454130 ref NP_006280.1 pTLN - talin gil3454130 ref NP_006280.1 pTLN - talin	841	80376318 (1681,	, 1682)		Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	264764
87002847 (1865, 1668) (Novel Protein sin. Galank gil382232gilgl8A44522.11 - Conflains protein domain (PF00170) - Struct 17241439 (1667, 1669) (Novel Protein sin. Galank gil2247210pjlAA20844] - Conflains protein domain (PF00186) - Instruction factor 18346844 (1869, 1869) Novel Protein sin. Galank gil2247210pjlAA20844] - Conflains protein domain (PF00186) - Instruction factor 18346844 (1869, 1869) Novel Protein sin. Galank gil22573pilA38299 - virulence Conflains protein domain (PF00180) - INCLASSIFIED Novel Protein sin. Galank gil22673pilA38299 - virulence Conflains protein domain (PF001020) - UNCLASSIFIED Novel Protein sin. Galank gil224721 Novel Protein sin. Galank gil224730jerjlA3829 Novel Protein sin. Galank gil224721	842	80078724 (1683,	, 1684)	-	Contains protein domain (PF00569) - Zinc finger present in dystrophin, CBP/p300	UNCLASSIFIED	264905, 264908, 265008, 265009, 18108374, 56182323, 264558
17941438 (1667, 1689) Nover Protein sim. GBank gigzzadzzi (bigilazzagia) 17041438 (1667, 1689) Nover Protein sim. GBank gigzzadzzi (bigilazzagia) Nover Protein sim. GBank gigzzadzi (bigilazzagia) Nover Protein s	843	87002847 (1685,	, 1686)		Contains protein domain (PF00170) - bZIP transcription factor	struct	264091, 29331825, 264906, 264768, 264563
1836644 (1889 1980) Nover Protein sim. GBank gijg28579[pir] A36829 - virulence Kinase T8863341 (1891 1892) Nover Protein sim. GBank gijg28579[pir] A36829 - virulence Contains protein domain (PF0120) - UNCLASSIFIED T8917849 (1893 1790) Nover Protein sim. GBank gijg2854 (1894 1895) Nover Protein sim. GBank gijg3138245199[Pig04] CASE) Contains protein domain (PF01608) - UNCLASSIFIED T10ASE) S5320333 (1701 1702) Nover Protein sim. GBank gijg44130[ref] NP_006280 IpT.IN - talin ULVEQ domain PF01608 T10ASE	844	17941439 (1687	, 1688)		Contains protein domain (PF00096) - Zinc finger, C2H2 type	transcriptfactor	265011
78693441 (1691, 1692) Novel Protein sun. GBank gilb22679jprijk35929 - virulence 78695346 (1693, 1694) 78617346 (1693, 1694) 78617346 (1693, 1694) 78617346 (1693, 1694) 78617346 (1693, 1694) 78617346 (1693, 1694) 78617346 (1693, 1694) 78617349 (1693, 1696) 78617349 (1693, 16	845	18346844 (1689,	1690)				264629
7883784 1893 1894 1895	846	79863441 (1691,	, 1692)	Novel Protein sim. GBank gilt regulatory protein VsrB - Psei		kinase	264907
78817469 (1869, 1869) 100CASSIFED 18756397 (1869, 1869) 100CASSIFED 18756397 (1869, 1869) 100CASSIFED 18756397 (1869, 1700) 100CASSIFED 18877469 (1869, 1700) 100CASSIFED 18877469 (1869, 1700) 100CASSIFED 163528 1700) 100CASSIFED 163528 1700) 100CASSIFED 163528 1700) 1702) 100CASSIFED 163528 1700) 1702) 17	847	78695348 (1693,	. 1694)	-		1,5	264909
19975307 (1897, 1898) 100CLASSIFIED 19917849 (1899, 1700) Novel Protein sim. GBank Contains protein domain (PF00120) - UNCLASSIFIED Glutamine synthetase CLUTAMINE SYNTHETASE (GLUTAMINE GUIDANINE	848	78489365 (1695,	, 1696)			UNCLASSIFIED	265020
78817849 (1659, 1700) Novel Protein sin. GBank	849	79756367 (1697,	, 1698)			UNCLASSIFIED	264566
Contains protein domain (PF01608) - Contains protein domain (PF01608) - Contains protein domain (PF01608) -	820	79817849 (1699,	. 1700)	JK_ECOLI - PUTATIVE	Contains protein domain (PF00120) - Glutamine synthetase	UNCLASSIFIED	264909
95320333 (1701, 1702) Novel Protein sim. GBank Contains protein domain (PF01608) - ULWEQ domain gil5454130[ref NP_006280.1pTLN - talin ULWEQ domain				IINE SYNIHEIASE (
gi 5454130 ref NP_005280.1 pTLN - talin	851	95320333 (1701,	1702)		Contains protein domain (PF01608) -		264488, 52644507, 264489, 18108398,
10147386 (1703 1704)				J.1pTLN - talin	I/LWEQ domain		65274572, 56182575, 22278994, 22278995,
10147366.1703 1704)			-				22278996, 35696286, 22278997, 22278998.
101473661703 1704)							22278999, 20281171, 264490, 264259,
10147366 1703 170a)			_				52645080, 29331822, 29331824, 66/1411/,
10147366 1703 1704)							23331023, 00432203, 23331020, 23331027, 35566052, 23331027, 35566052, 23331027, 3676408, 36146409
10147366 (1703 1704)							264107 264905 264906 264907 264908
1014736 L1703 1704)							52844045 56182435 265006 265007
10147366 11703 1704)							265008, 265009, 264910, 60432229.
10147366 11703 17041							60431735, 60433356, 33657402, 60433438,
1014736 1703 1704		-					264595, 264758, 264759, 21906754,
10147366 11703 17041							33109954, 52644296, 265010, 265011,
10147366 1703 17041							87168559, 265017, 265018, 265019, 264760,
1014736 1703 1704)							264761, 264762, 264681, 18108351, 264763,
1014736 1703 17041		_					264448, 264682, 264764, 264683, 18108354,
1014736 1703 1704)							264288, 264369, 264685, 264766, 264687.
1014736 11703 1704)							264768, 264769, 21906765, 21906766,
1014736 11703 1704)							21906767, 21906768, 29148627, 21906769,
1014736 1703 1704)	_						29148629, 55811957, 35695917, 265020,
1014736 11703 1704)						-	265021, 265022, 60170615, 52644150.
1014736 11703 1704)							264691, 264692, 33657023, 264693, 263966.
10147366 11703 1704)		-					33657109, 27486261, 27486262, 27486264,
10147366 11703 1704)							27486265, 35695763, 60431602, 18108370,
10147366 (1703-1704)							20281069, 264629, 18108374, 18108376,
10147366 (1703-1704)							55811576, 35696423, 35695855, 264634,
	862	10147366 /1703	1704)				264635, 264636, 264555, 60431850, 264556,

650	142022607 /470E 470E	470E 470	-			
250	13032301	1707 470	in the Court of th			264636
3	9643690	70,1	(V16136) 2-enoate reduction (M16136) 2-enoate reductions (Moorella thermoacetica)		reductase	264568
855	79641130 (1709, 1710)	1709, 171	(0)			264692
856	11594236 (1711, 1712)	1711, 171	(2)		UNCLASSIFIED	264591
857	79210165 (1713, 1714)	1713, 171	(4)		UNCLASSIFIED	264630, 264634
858	80248910 (1715, 1716)	1715, 171	(9)			265008, 265009, 264601, 264602, 264603, 18108351
859	20296634 (1717, 1718)	1717, 171	(8)			264559
860	80041749 (1719, 1720)	1719, 172	(0;		UNCLASSIFIED	264489
861	65857045 (1721, 172	(2)		UNCLASSIFIED	33657023, 264630
862	80079467 (1723, 1724)	1723, 172	(4)			264600
863	80579931 (1725, 172	80579931 (1725, 1726) Novel Protein sim. GBank gi 2246532 (U93872) - ORF 73.		UNCLASSIFIED	264488, 18108398, 35696286, 264259.
			contains large complex repeat CR 73 [Kaposi's sarcoma-			18108351, 264288, 265021
٦			associated herpesvirus			
864	94939904 (1727. 172		_	UNCLASSIFIED	264259, 264112, 263974
	80045310 (1729, 173	5689884 emb CAB52047.1 - ein [Streptomyces coelicolor	Contains protein domain (PF01479) - S4 domain		264635, 264600, 264636, 264591, 264602, 264693
930	, +6000	4724				
8) [6020] (8	3/1,18/1	60102031 (1731, 1732) Nover Protein sim. GBank gil455786/refl/P_000341.1 pABCR - ATP binding cassette		transport	264288, 264557, 264558
\int			transporter			
867	80062402 (1733, 1734)	1733, 173	(4)			264605
868	10075364 (1735, 1736)	1735, 173	(9)		UNCLASSIFIED	564909
698	80062406 (1737, 1738)	1737, 173				264605, 264687, 18108374
870	80249651 (1739, 174	80249651 (1739, 1740) Novel Protein sim. GBank gij628660 pir S37755 - Adenylyl- transferase - Escherichia coli		transferase	264601, 264636
	, 2000000					
8/1	, 68287602	1/41, 1/4	ZU378295 (1741, 1742) Novel Protein sim. GBank gij1708180jspjQ10602jHEMK_MYCTU - HEMK PROTEIN HOMOLOG		UNCLASSIFIED	264603
872	95197114 (1743, 174	95197114 (1743, 1744) Novel Protein sim. GBank gij 1545959 emb CAA67763 -		UNCLASSIFIED	35696286, 22278998, 264259, 29331822.
			(X99384) paladin [Mus musculus]			29331824, 29331825, 60432289, 29331826, 29331827, 29331828, 35696052, 264509, 264906, 264907, 264908, 264909, 264906, 264907, 264908, 264909, 265008, 265008, 264910, 264591, 60433356, 264506, 264603, 264601, 265018, 264605, 264602, 264603, 265017, 265018, 264605, 264629, 264629, 264629, 264634, 264631, 264541, 264531, 264638, 264639, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 262739002, 22279000, 22279000,
						264564
873	20189728 (1745, 174	20189728 (1745, 1746) Novel Protein sim. GBank gild196104 (AE001569) - putative C Outer membrane protein [Helicobacter pylori J99]	4156104 (AE001569) - putative Contains protein domain (PF00096) - UNCLASSIFIED licobacter pylori J99) Zinc finger, C2H2 type	UNCLASSIFIED	264595

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6 6	60077892 (1747, 1748)	8UU/1592 (1/4/, 1/46) NOVER TOLEIN SIM. GBARK gil13419sp[PORB9]SCRB_BACSU - SUCROSE-6- PHOSPHATE HYDROL ASF (SICRASE) (INVERTASE)		UNCLASSIFIED	264600
875	86608446 (1749, 1750)				264259, 264448, 264288, 264557, 87168518
876	86465157 (1751, 1752)	86465157 (1751, 1752) Novel Protein sim. GBank gil3128283 (AF010496) - iron(iii) dicitrate transport ATP-binding protein [Rhodobacter capsulatus]	Contains protein domain (PF00005) - transport ABC transporter	transport	<u>264907, 264601, 264602, 264605, 265020,</u> 60431602
877	87802548 (1753, 1754)	87802548 (1753, 1754) Novel Protein sim. GBank gij731074 sp P40349 URB1_USTMA - SIDEROPHORE BIOSYNTHESIS REGULATORY PROTEIN URBS1	Contains protein domain (PF00320) - transcriptfactor GATA zinc finger	transcriptfactor	22278998, 264909, 264369
878	80187289 (1755, 1756)	80187289 (1755, 1756) Novel Protein sim. GBank gil1351614[sp]Q09853[YAED_SCHPO - HYPOTHETICAL 181.5 KD PROTEIN C23D3.13C IN CHROMOSOME I		ATPase_associated 264369, 264555	264369, 264555
878	94328962 (1757, 1758)	94328962 (1757, 1758) Novel Protein sim. GBank gij3875304jembjCAA98434j - (Z74030) predicted using Genefinder; cDNA EST EMBL:C07609 comes from this gene; cDNA EST EMBL:C09023 comes from this gene; cDNA EST yk489h9.3 comes from this gene; cDNA EST yk489h9.3 comes from this gene; cDNA EST yk489h9.5 comes from			56182575, 29331824, 264508, 264906, 265018, 18108351, 264448, 264683, 21906766, 21906768, 60170615, 33657023, 65274620, 33657109, 18108374, 35695855, 264563
880	8491135 (1759, 1760)	Novel Protein sim. GBank gil137120 sp P11214 UROT_MOUSE - TISSUE PLASMINOGEN ACTIVATOR PRECURSOR (TPA) (T- PLASMINOGEN ACTIVATOR)	Contains protein domain (PF00051) - cathepsin Kringte domain	cathepsin	264508
881	11290122 (1761, 1762)			UNCLASSIFIED	264508
882	11077011 (1763, 1764)	11077011 (1763, 1764) Novel Protein sim. GBank gil 2632098 emb CAA75667 - (Y15513) Prodos protein [Drosophila melanogaster]		UNCLASSIFIED	264558
883	79582969 (1765, 1766)			UNCLASSIFIED	264688
884	13517921 (1767, 1768)	13517921 (1767, 1768) Novel Protein sim. GBank gi 1155068 emb CAA64425 - (X94976) cell wall-plasma membrane linker protein [Brassica napus]		UNCLASSIFIED	264636
885	80052457 (1769, 1770)	80052457 (1769, 1770) Novel Protein sim. GBank gi 2078027 emb CAB08467 - (Z95208) hypothetical protein Rv2372c [Mycobacterium [tuberculosis]		UNCLASSIFIED	264605, 18108362
886	(11685136 (1771, 1772)				264690
887	94315307 (1773, 1774)	94315307 (1773, 1774) Novel Protein sim. GBank gil2695834 emblCAA15904 - (AL021008) sucA (Mycobaclerium tuberculosis)		dehydrogenase	35696052, 264906, 264600, 264603, 35695917, 35695855, 264636
888	10083399 (1775, 1776)	10083399 (1775, 1776) Novel Protein sim. GBank gil5689395 db BAAB2981.1 - (AB028952) KIAA1029 protein [Homo sapiens]		UNCLASSIFIED	264908
888	20385917 (1777, 1778)	20385917 (1777, 1778) Novel Protein sim. GBank gi[1881338 dbj BAA19365 - (AB001488) PROBABLE INTEGRAL MEMBRANE PROTEIN, SIMILAR TO CHLORAMPHENICOL RESISTANCE PROTEIN OF STREPTOMYCES VENEZUELAE. [Bacillus subiilis]		:	264603
890	19904337 (1779, 1780)	19904337 (1779, 1780) Novel Protein sim. GBank gij854065[emb CAA58337] - (X83413) U88 [Human herpesvirus 6]			264629

				Γ				Г				ni ni			<u> </u>	Τ	Г		
264636	22278996, 22278999, 29331828, 35696052, 264908, 264909, 265009, 265011, 264602, 265019, 264766, 21906765, 21906766, 21906769, 265020, 265021, 56526486	264689, 263967	263978	264508, 264600, 264555, 264559	264632	264686, 29331828, 264511	56182575, 60432289, 56182435, 60432229, 55811957, 22279000, 264486	264601	264769, 264691, 264563	264907, 264602, 264605, 264769, 35695917, 18108376, 264563	264909, 265008, 264602, 264604, 264769, 264689, 264693	35696052, 264905, 264510, 264511, 264612, 264605, 264760, 18108351, 264762, 264687, 264768, 254769, 264688, 21906764, 35695917, 27486262, 35695855, 264634, 264636, 264486	264604	264909	264595, 264605	60432289, 264601, 264690	264638	264602	265009
UNCLASSIFIED	transcriptfactor	synthase		UNCLASSIFIED		UNCLASSIFIED			transport	synthase	isomerase	kinase	hydrolase	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	hda
						Contains protein domain (PF00047) - UNCLASSIFIED Immunoglobulin domain				Contains protein domain (PF01411) - synthase IRNA synthetases class II (A)	Contains protein domain (PF00330) - isomerase Aconitase family (aconitate hydratase)	Contains protein domain (PF00370) - kinase FGCY family of carbohydrate kinases	Contains protein domain (PF00491) - hydrolase Arginase family						Contains protein domain (PF00684) - eph
13516879 (1781, 1782) Novel Protein sim. GBank gil4959396[gb]AAD34331.1 AF11248 - (AF112481) RAD54B protein Homo saniens1	87634157 (1783, 1784) Novel Protein sim. GBank gij545526jbbs 143833 - LBP- 1b=transcription factor binding to initiation site of HIV-1 (alternatively spliced) [human, Namahva cells, Peptide, 541 aa]	79168037 (1785, 1786) Novel Protein sim. GBank gi 2829688 sp P80608 CYSK_MAIZE - CYSTEINE SYNTHASE (O-ACETYLSERINE SULFHYDRYLASE) (O- ACETYLSERINE (THIOL)-LYASE) (CSASE)	-						80499768 (1799, 1800) Novel Protein sim. GBank gi 1750127 (U66480) - YncC Bacillus subtilis	80502410 (1801, 1802) Novel Protein sim. GBank gij3122879 sp O07438 SYA_MYCTU - ALANYL-TRNA SYNTHETASE (ALANINE-TRNA LIGASE) (ALARS)	80503301 (1803, 1804) Novel Protein sim. GBank gij3355701 embjCAA20001 - (AL031124) 3-isopropylmalate dehydratase large subunit [Streptomyces coelicolor]	82060208 (1805, 1806) Novel Protein sim. GBank gi 2960120 emb CAA18018.1 - (AL022121) glpK [Mycobacterium tuberculosis]	20451078 (1807, 1808) Novel Protein sim. GBank gi 728887 sp P40906 ARG _COCIM - ARGINASE	Novel Protein sim. GBank gil4567200[gb AAD23616.1 AC00716 - (AC007168) hypothetical protein [Arabidopsis thaliana]	80052628 (1811, 1812) Novel Protein sim. GBank gij3738200 emb CAA21292 - (AL031855) pulative vacuolar membrane protein Schizosaccharomyces sombel	_		20727907 (1817, 1818) Novel Protein sim. GBank gij3868940 db BAA34296 - /AB015054) Alq2 [Rhizomucor pusillus]	16776206 (1819, 1820) Novel Protein sim. GBank gil4589726 dbj BAA76883.1 -
, 1782)	, 1784)	, 1786)	, 1788)	(1790)	(1792)	1794)	, 1796)	(1798)	. 1800)	, 1802)	, 1804)	, 1806)	. 1808)		, 1812)	1814)	1816)	, 1818)	, 1820)
79 (1781	57 (1783	37 (1785	11102240 (1787, 1788)	80239868 (1789, 1790)	79747803 (1791, 1792)	23 (1793	87895109 (1795, 1796)	11100463 (1797, 1798)	68 (1799	10 (1801	01 (1803	08 (1805	78 (1807	9398483 (1809, 1810)	28 (1811	87913201 (1813, 1814)	11754482 (1815, 1816)	07 (1817	06 (1819
135168	876341	791680.	111022	802398	797478	949919.	878951	111004	804997	805024	805033	820602	204510	939848	800526	879132	117544	207279	167762
891	892	893	8 9	895	968	897	898	668	006	901	805	903	904	902	906	206	908	606	910

91	87454340 (1821, 1822)	87454340 (1821, 1822) Novel Protein sim. GBank gi 548774 sp P35685 RL7A_ORYSA - 60S RIBOSOMAL PROTEIN L7A		ribosomalprot	265010, 264604, 60432113
912	20448863 (1823, 1824)	20448863 (1823, 1824) Novel Protein sim. GBank gil2314008 gb AAD07921.1 - (AE000597) CDP-diglyceride hydrolase (cdh) {Helicobacter pylori 26695]		hydrolase	264559
913	20469357 (1825, 1826)			UNCLASSIFIED	264604
914	79183351 (1827, 1828)	79183351 (1827, 1828) Novel Protein sim. GBank gij417657 sp Q03604 RIR1_CAEEL - PROBABLE RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE LARGE CHAIN (RIBONUCLEOTIDE REDUCTASE)	Contains protein domain (PF00317) - reductase Ribonucleotide reductase	reductase	264636
915	87606703 (1829, 1830)	87606703 (1829, 1830) Novel Protein sim. GBank gij5689571 dbj BAA83069.1 - (AB029040) KIAA1117 protein [Homo sapiens]			18108398, 22278996, 66714117, 264906, 264591, 21906768, 265020, 55811576, 264638
916	79444091 (1831, 1832)	79444091 (1831, 1832) Novel Protein sim. GBank gil4186110 emb CAA71790 - (Y10831) putative integrase [Ralstonia eutropha]		UNCLASSIFIED	264595
917	20195985 (1833, 1834)			UNCLASSIFIED	264605
918	91226795 (1835, 1836)	91226795 (1835, 1836) Novel Protein sim. GBank gil1655699 emb CAA69032 - (Y07752) pherophorin-S [Volvox carteri]		synthase	264259, 29331826, 264908, 265019, 264448, 265020, 264635, 83373044
919	80436785 (1837, 1838)	80436785 (1837, 1838) Novel Protein sim. GBank gil5689968 emb CAB52005.1 - (AL.10963) putative membrane protein [Streptomyces coelicotor A3(2)]			265006, 264512, 264600, 264602, 264604, 264768, 18108370, 264563
920	79606095 (1839, 1840)	79606095 (1639, 1840) Novel Protein sim. GBank gij1168448jspjQ05813JAMP1_STRLI - XAA-PRO AMINOPEPTIDASE I (X-PRO AMINOPEPTIDASE I) (AMINOPEPTIDASE P I) (APP) (PEPP I) (AMINOACYLPROLINE AMINOPEPTIDASE I)		peplidase	264508
921	19858634 (1841, 1842)	19858634 (1841, 1842) Novel Protein sim. GBank gil3850084 emb CAA21911.1j - (AL033389) alcohol dehydrogenase [Schizosaccharomyces pombe]		UNCLASSIFIED	264600
922	78982605 (1843, 1844)				265019, 22279002
923	86695830 (1845, 1846)	86895830 (1845, 1846) Novel Protein sim. GBank gil267079 sp P29514 TBB6_ARATH - TUBULIN BETA-6 CHAIN		tubulin	264907, 265008, 265009, 265010, 18108351, 264689, 265021, 18108370, 18108374, 18108385
924	21431341 (1847, 1848)				264510
925	20630332 (1849, 1850)	20630332 (1849, 1850) Novel Protein sim. GBank		esterase	264603
		gi[2497688]spiQ60963]PAFA_MOUSE - PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE PRECURSOR (PAF ACETYLHYDROLASE) (PAF 2-ACYLHYDROLASE) (LDL-ASSOCIATED PHOSPHOLIPASE A2) (LDL-PLA(2)) (2-ACETYL-1-ALKYLG-YCEROPHOSPHOCHOLINE ESTERASE) (1-ALKYL-2-ACETYLGLYCEROPHO			
926	79397657 (1851, 1852)	79397657 (1851, 1852) Novel Protein sim. GBank gil3882325 dbj BAA34522.1 - (AB018345) KIAA0802 protein [Homo sapiens]			558 11957, 263972, 264639
927	37036201 (1853, 1854)			UNCLASSIFIED	264769

80070610 (1855, 1856)	80070610 (1855, 1856) Novel Protein sim. GBank gi[156146 (M30316) - xanthine dehydrogenase (AA at 2538) [Calliphora vicina]		dehydrogenase	264605
-+			UNCLASSIFIED	264603
_ =	Novel Protein sim. GBank giļ4115936jgbļAAD03446.1Į - (AF118223) No definition line found [Arabidopsis thaliana]			264259
	10245731 (1861, 1862) Novel Protein sim. GBank gil4490609 emb CAB38642.1 - (AJ133495) ribonucelotide reductase major subunit (Staphylococcus aureus)	Contains protein domain (PF00317) - reductase Ribonucleotide reductase	reductase	264486
	Novel Protein sim. GBank gi[5459396 emb CAB50754.1 - (AL096839) putative integral membrane transport protein [Streptomyces coelicolor]		transport	264508, 264905, 264906, 264909, 264600. 264602, 264603, 264605, 264766, 264558, 18108387, 264486
	94326010 (1865, 1866) Novel Protein sim. GBank gil5689523 dbj BAA83045.1 - (AB029016) KIAA1093 protein [Homo sapiens]	Contains protein domain (PF00627) - UNCLASSIFIED UBA domain	UNCLASSIFIED	264508, 264686, 264693, 27486261, 18108370, 65274791, 264636, 264559,
	80039105 (1867, 1868) Novel Protein sim. GBank gi 119111 sp P12978 EBN2_EBV - EBNA-2 NUCLEAR PROTEIN			264369, 22279002
80063162 (1869, 1870)			UNCLASSIFIED	264604, 264605, 264693, 18108370, 18108374
	80026632 (1871, 1872) Novel Protein sim. GBank gi[845686 (M32103) - ORF-27 [Staphylococcus aureus]		UNCLASSIFIED	22278995, 22278996, 264602, 264687, 32833986, 18108387
	80250273 (1873, 1874) Novel Protein sim. GBank gi 1360669 pir CGHU1V - Collagen alpha 1(V) chain precursor - human		synthase	263978
_ T	80026633 (1875, 1876) Novel Protein sim. GBank gi 2226243 emb CAA74531.1 - (Y14083) hypothetical protein [Bacillus subtilis]		UNCLASSIFIED	264602
110/1694 (1877, 1878)				264600
	94144252 (1879, 1880) Novel Protein sim. GBank gi]3560166[emb CAA20678] - (AL031525) ubiquitin carboxyt-terminal hydrolase [Schizosaccharomyces pombe]	Contains protein domain (PF00442) - ubiquitin Ubiquitin carboxyl-terminal hydrolases family 2	ubiquitin	264905, 264906, 264907, 264908, 264909, 264511, 264910, 264592, 33657402, 264596, 264760, 264683, 264766, 264768
				264769, 33657023, 33657109, 264628, 264629, 264630, 264635, 264636, 264555,
				264637, 264556, 264638, 264639, 83373044, 18108385, 264565, 18108391
11398414 (1881, 1882)			Г	264593
13404122 (1883, 1884)	13404122 (1883, 1884)		UNCLASSIFIED	264760
~	oucousso (1965), 1960) Nover Protein Sim. GBank gil4033729 (AF-038595) - apolipoprotein N-acyltransferase [Pseudomonas aeruginosa]		UNCLASSIFIED	264600, 264687, 264689, 264563
<u> </u>	Novel Protein sim. GBank gijz494764[sp]Q50729]GUAA_MYCTU - GMP SYNTHASE (GLUTAMINE-HYDROLYZING) (GLUTAMINE AMIDOTRANSFERASE) (GMP SYNTHETASE)		synthase	264511, 264603
	60032477 (1889, 1890) Novel Protein sim. GBank gil732353 sp P39606 YWCH_BACSU - HYPOTHETICAL 36.6 KD PROTEIN IN QOXD-VPR INTERGENIC REGION	Contains protein domain (PF00818) - UNCLASSIFIED Ice nucleation protein repeat	UNCLASSIFIED	264906, 264604, 264605, 265020, 18108387
79248402 (1891, 1892)				265017

947	81802699 (1893, 1894)		Contains protein domain (PF00459) - phosphatase Incsitol monophosphatase family	phosphatase	18108394, 22278996, 264907, 264909, 265008, 265009, 264000, 264000, 264758, 264689, 264689, 264689, 264689
948	88165538 (1895, 1896)	88165538 (1895, 1896) Novel Protein sim. GBank gi 2827284 (AF041037) - novel antagonist of FGF signaling [Homo sapiens]		lg/	18108398, 56182575, 22278997, 22278999, 60432049, 29331822, 29331826, 264907, 56182435, 55811386, 265011, 264600, 265017, 265018, 265019, 18108351, 265020, 265021, 265022, 27486265, 263972, 5811576, 264638, 60170394, 264566
948	88081786 (1897, 1898)	88081786 (1897, 1898) Novel Protein sim. GBank gil4507985 ref NP_003427.1 pZNF1 - zinc finger protein 135 Zinc finger, C2H2 type (clone pHZ-17)	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	29331825, 21906764, 27486261, 21906766, 52644296, 33657349, 87168518, 56994075, 265020, 265021, 87168559, 52644150, 264637
920	79485872 (1899, 1900)	79485672 (1899, 1900) Novel Protein sim. GBank gi 1079461 pir S43865 - cytokeratin 8, type II - potoroo (fragment)	Contains protein domain (PF00038) - struct intermediate filament proteins	struct	264683, 18108361
951	20451411 (1901, 1902)	20451411 (1901, 1902) Novel Protein sim. GBank gil5420387 emb CAB46679.1 - (AJ243459) proteophosphogiycan [Leishmania major]		UNCLASSIFIED	264604
952	79566954 (1903, 1904)	- ≏ থ		cathepsin	264910, 264691
953	10196003 (1905, 1906)	10196003 (1905, 1906) Novel Protein sim. GBank gi[2495642[sp]Q47142[yFHS_ECOLI - HYPOTHETICAL 41.6 KD PROTEIN IN CSIE-GLYA INTERGENIC REGION		transport -	264510
\$\$ \$	9893326 (1907, 1908)	Novel Protein sim. GBank gij2360965 (AF016253) - D- amino acid dehydrogenase [Klebsiella aerogenes]		dehydrogenase	264508
955	95313410 (1909, 1910)	95313410 (1909, 1910) Novel Protein sim. GBank gil5454064 ref NP_006319.1 pSIP - SYT interacting protein	Contains protein domain (PF00076) - IpSIPI - SYT interacting protein RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	dna_rna_bind	56994075, 264509, 264905, 264906, 264907, 264908, 264909, 264910, 264910, 264758, 264759, 264760, 264910, 264758, 264762, 264763, 264764, 264766, 264686, 264767, 264768, 264769, 26469, 264687, 264630, 264631, 264632, 264634, 264635, 264636, 264631, 264638, 264634, 264639, 18108388, 264563, 264564
926	80064224 (1911, 1912)	80064224 (1911, 1912) Novel Protein sim. GBank gi 2052129 emb CAB08155 - (294752) rimJ [Mycobacterium tubercutosis]			264605
957	80056206 (1913, 1914)	80056206 (1913, 1914)		UNCLASSIFIED	264603, 18108362
928	80036446 (1915, 1916)	i.		UNCLASSIFIED	264908, 264910, 264762, 263978, 264637
656	80026647 (1917, 1918)	80026647 (1917, 1918) Novel Protein sim. GBank gil2131050 emb CAB09260 - (295844) opcA [Mycobacterium tuberculosis]		UNCLASSIFIED	264602, 264692
096	37815406 (1919, 1920)	37815406 (1919, 1920) Novel Protein sim. GBank gil2129478 pir S51939 - chitinase (EC 3.2.1.14) precursor - beet		ONCLASSIFIED	264259
961	20567383 (1921, 1922)	-			263978
962	11399318 (1923, 1924)				264593

963	80590374 (1925, 1926)	1926)			UNCLASSIFIED	264510, 264288, 264555, 264556, 264559, 264486
964	79832019 (1927, 1	1928)	79832019 (1927, 1928) Novel Protein sim. GBank gil4589622[dbj BAA76833.1 - (AB023206) KIAA0989 protein [Homo sapiens]		UNCLASSIFIED	264112, 264910, 264689
596	91229485 (1929, 1	1930)	Novel Protein sim. GBank gi 5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	264488, 265017, 264448, 264634, 264558, 83373044
996	95292815 (1931, 1932)	1932)			UNCLASSIFIED	264906, 264592, 264596, 264604, 264768, 21906764, 264692, 284693, 264629, 264636, 264638
967	79255708 (1933, 1	1934)	79255708 (1933, 1934) Novel Protein sim. GBank gil1731207[sp]Q11156 RGX3_MYCTU - SENSORY TRANSDUCTION PROTEIN REGX3	Contains protein domain (PF00072) - Response regulator receiver domain	phosphatase	264760
896	79560269 (1935, 1	1936)	79560269 (1935, 1936) Novel Protein sim. GBank gil2651836 emb CAA75187 - (Y14964) putative transport protein [Methylophilus methylotrophus]		transport	264693
696	79919470 (1937, 1	1938)	79919470 (1937, 1938) Novel Protein sim. GBank gi[5419878 emb CAB46422.1 - (AL096747) hypothetical protein [Homo sapiens]	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type	dna_ma_bind	35696286, 264685, 264686, 35695917, 264692, 18108374, 264635
026	95085947 (1939, 1940)	1940)			UNCLASSIFIED	18108392, 18108394, 18108398, 22278995, 22278996, 22278998, 22278999, 29147620, 264828, 265006, 265007, 265008, 265009,
						18108348, 33109954, 265010, 265011, 18108351, 264288, 21906767, 21906768, 18108370, 18108377, 264630, 264635, 18108380, 83373044, 18108387, 18108388
971	78919770 (1941, 1942)	1942)			UNCLASSIFIED	265007, 265020, 22279002
972	20710704 (1943, 1944)	1944)				264557
973	20370183 (1945, 1	1946)	20370183 (1945, 1946) Novel Protein sim. GBank gi 1723119 sp P53990 Y174_HUMAN - HYPOTHETICAL PROTEIN KIAA0174			264604
974	80057103 (1947, 1948)				UNCLASSIFIED	264565
975	10196018 (1949, 1	1950)			UNCLASSIFIED	264510
976	80205742 (1951, 1	1952)	80205742 (1951, 1952) Novel Protein sim. GBank gij3881459jembjCAA92988.1]		UNCLASSIFIED	264508, 264906, 264758, 264632, 264639,
			(CD6733) predicted using Genelinder; Similarity to Yeast hypothetical protein YIK9 (SW:YIK9_YEAST); cDNA EST EMBL:D27680 comes from this gene; cDNA EST EMBL:D27679 comes from this gene; cDNA EST EMBL:D64477 comes from this ge			264.56.3
977	10355349 (1953, 1	1954)			UNCLASSIFIED	264906
978	80025927 (1955, 1956)				UNCLASSIFIED	264600, 264602, 264603, 264604
979	80447820 (1957, 1	1958)	80447820 (1957, 1958) Novel Protein sim. GBank gil3171904 emb CAA75869 - (Y15908) DIA-12C protein [Homo sapiens]		UNCLASSIFIED	264767, 264768, 265006, 265007, 264906
980	80025928 (1959, 1	1960)	80025928 (1959, 1960)		UNCLASSIFIED	264600, 264602, 264605
981	80098550 (1961, 1	1962)	Novel Protein sim. GBank gij3599940 (AF017368) - faciogenital dysplasia protein 2 [Mus musculus]		UNCLASSIFIED	264692, 264555, 264556, 264557, 264559

17.102610 (1962, 1964) Word Protein Sim. GBank g 23.23297 AEON 1963 (11981, 1992) Word Protein Sim. GBank g 23.17179 (1989, 1990) Word Protein Sim. GBank g 23.17179 (40.004227) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004227) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004227) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.00427) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Contains protein domain (PF00176) Word Protein Sim. GBank g 23.17179 (40.004027) Word Protein Sim. GBank g 23.17179 (40.004027) Word Protein	600	105670	1062 106	Maint Bratala sim CBank allocknoonlambiC 4471478		INCLASSIFIED	264404
100969504 1965, 1969 Navel Protein am. CBank gild31504jemb[CA407660 1] - Conlains protein domain (PF00534) - Lansfersse (5461386 (1965, 1970) Navel Protein sim. CBank gild31504jemb[CA407660 1] - Conlains protein domain (PF00534) - Lansfersse (5461386 (1969, 1970) Navel Protein sim. CBank gild31504jemb[CA407660 1] - Conlains protein domain (PF00534) - Lansfersse (1976, 1972) Conlains protein domain (PF00534) - Lansfersse (1976, 1972) Conlains protein domain (PF00537) Lansfersse (1976, 1972) Conlains protein domain (PF01421) - Conlains protein domain (PF007121) - Conlains (CLCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC		0.0008109	1903, 190	(Y10545) fused-ccdB (Escherichia coli)			
100 100		90995041 (1965, 196	Novel Protein sim. GBank gil476389 pir B43402 - myosin		struct	65274572, 56182575, 264908, 264909, 265007 265008, 264758, 265010, 55811150.
UNCLASSIFED 1982 1997 Novel Protein sim GBank gi[245;504 emb[CAA07660.1] - Contains protein domain (PF0033) - Itansferase UNCLASSIFED 19805239 (1971; 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1974 1972 1975 1				meavy Crain-D, heuronal - Chicker			33657023, 264634, 264557, 264558
564611956 (1986) 1970 Novel Protein ain. CBank gil44150lemPlCAA07660 11 - Contains protein domain (PF00534) - Iransferase group 1	Г	20466876 (1967, 196			UNCLASSIFIED	264605
80053247 (1983, 1984) 1982 1982 1982 1982 1982 1983, 1984 1982 1983 1983 1983 1983 1983 1983 1983 1984 1982 1983 1984 1982 1983 1984 1982 1984 1982 1984 1982 1984 1982 1984 1985 1984 1985 1984 1985 1984 1985		65461368 ((1969, 197		Contains protein domain (PF00534) - Glycosyl transferases group 1	transferase	56182435, 264600
1985/223 (1973 1974) 1985/2505(1975 1978) 1985/2505(1975 1978) 1985/2505(1975 1978) 1985/2505(1975 1978) 1985/2505(1977 1978) 1980/2505(1976 1978)	Г	87102868 (1971, 197.	-		UNCLASSIFIED	264106, 264110, 265020, 60170615
1986/566 (1977, 1976) 1986/566 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1977, 1976) 1986/5746 (1987, 1986) 1986/5746 (1987, 1987, 1987) 1987/5746 (1987, 1987, 1987) 1987/5746 (1987, 1987, 1984) 1982/5746 (1987, 1987, 1984) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987) 1982/5746 (1987, 1987, 1987, 1987) 1982/5746 (1987, 198	Г	79867231 (1973, 197	(†)			264909
80057746 (1977 : 1978) 80057746 (1977 : 1978) 80057746 (1979 : 1980) Novel Protein sim. GBank gil2302991 (AC004227) - Gontains protein domain (PF00421) - Oxidase gil572506igglyAAD40000 (1AF00015 · (AF000152) METH1 Reproyale (M172B) family zinc gil510553FiED protein lim. GBank gil2105049jemb CA808835 - Adenyale and Guanyiate cyclase luberculosis 11090590 (1987 : 1988) Novel Protein sim. GBank gil2329297 (AE001355) - Reductase luberculosis 11090590 (1987 : 1988) Novel Protein sim. GBank gil2329297 (AE001355) - Reductase luberculosis 11090590 (1987 : 1989) Novel Protein sim. GBank gil2317119 (AC004227) - Gontains protein domain (PF00059) - Itansport larsconder, ATP-binding protein larchaedglobus fulgidus ABC transporter larsconder, ATP-binding protein larchaedglobus fulgidus ABC transporter GLYCOPROTEIN PRECURSOR	Γ	19858661 (1975, 1970	(9)		UNCLASSIFIED	264600
89057746 (1979, 1980) Novel Protein sim. GBank gi[20105049] 101061407746 (1981, 1982) Novel Protein sim. GBank gi[20105049] 101061407745 (1991, 1992) Novel Protein sim. GBank gi[20105049] 1010614053 (1986, 1986) Novel Protein sim. GBank gi[20105049] 1010614053 (1987, 1980) Novel Protein sim. GBank gi[20105049] 1010614053 (1987, 1980) Novel Protein sim. GBank gi[20105049] 1010614053 (1987, 1980) Novel Protein sim. GBank gi[20105049] 10105050 (1987, 1980) Novel Protein sim. GBank gi[20107179 (AE001050] Contains protein domain (PF00076) - helicase gi[5105072]ppAAD33760 (1987, 1989) Novel Protein sim. GBank gi[20107179 (AE001001) - ABC Contains protein domain (PF00059) - phosphatase gi[5105072]ppAD33760 (1987, 1999) Novel Protein sim. GBank gi[20107179 (AE001001) - ABC Contains protein domain (PF00059) - transport Lansporter, ATP-binding protein sim. GBank gi[2649101 (AE001001) - ABC Contains protein domain (PF00005) - transport CONTROPROTEIN PRECURSOR	Т	88095329 (1977, 197	(8)		UNCLASSIFIED	264508, 265017, 264534, 264564
10106140 (1981, 1982) Protein Tronn Sapleria Internation Protein Sim. CBank gil2105049jemb(CAB08835] Contains protein domain (PF00211) - UNCLASSIFIED (205436) hypothetical protein R. CBank gil3329297 (AE001355) - Reductase (1987, 1986, 1986) Movel Protein sim. CBank gil2329297 (AE001355) - Reductase (10914053 (1987, 1980) Novel Protein sim. CBank gil2317179 (AC004227) - Contains protein domain (PF00176) - Intersport (1993, 1994) Novel Protein sim. CBank gil2611719 (AC004227) - Reductase (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Reductase (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Reductase (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Reductase (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein Sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein Sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein Sim. CBank gil2611719 (AC004227) - Red (1397390 (1995, 1999) Novel Protein Sim. CBank gil2611717		88057746 ((1979, 198	1/AF06015 - (AF060152) METH1		oxidase	.264259, 264908, 265009, 264910, 264596, .264369, 264288, 264766, 264628, 264635, 264668
109614053 (1981, 1982) 100	T			_			00000
10814053 1984 Novel Protein sim. GBank gil29105049lemb CAB08835 - Contains protein domain (PF00213) UNCLASSIFIED (295436) hypothetical protein Rv3645 [Mycobacterium Adenylate and Guanylate cyclase Iuberculosis 10814053 (1985, 1986) Novel Protein sim. GBank gil29297 (AE001355) - Ranscriptonal adrivator SRCAP [Homo sapiens] SNF2 and others N-ferminal domain (PF00176) - Italy phosphalase (1993, 1994) Novel Protein sim. GBank gil2911719 (AC004227) - PDZ domain (Also known as DHR or (I1397390 (1995, 1995) Novel Protein sim. GBank gil2649101 (AE001001) - ABC Contains protein domain (PF00055) - Itansport (I1397390 (1995, 1996) Novel Protein sim. GBank gil2649101 (AE001001) - ABC Contains protein domain (PF00055) - Itansport (I1397390 (1995, 1996) Novel Protein sim. GBank gil2649101 (AE001001) - ABC Contains protein domain (PF00055) - Itansport (I1397390 (1995, 1996) Novel Protein sim. GBank gil2649101 (AE001001) - ABC Contains protein domain (PF00055) - Itansport (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH (I1397390 (1995, 1996) Novel Protein sim. GBank gil2040 - HISTIDINE-RICH	٦	10106140 (1981, 198	2)		UNCLASSIFIED	264909
110914053 (1985, 1986) Novel Protein sim. GBank gi]3329297 (AE001355) - Rabonalis Rabonacide Reductase, Large Chain [Chlamydia SNF2 and others N-terminal domain (PF00176) - helicase gils 106572[glp/AAD39760.14F14394 - (AF143946) SNF2 and others N-terminal domain (PF00176) - helicase gils 106572[glp/AAD39760.14F14394 - (AF143946) SNF2 and others N-terminal domain (PF00176) - helicase gils 106572[glp/AAD39760.14F14394 - (AF143946) AEC Contains protein domain (PF00176) - transport (ARA001LB [Homo sapiens] PDZ domain (Also known as DHR or GLGF) - transport (ARA001LB [Homo sapiens] GLGF) ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (Contains protein domain (PF00005) - transport (ARA001LB [Homo sapiens] ABC (ARA001LB (ARA001LB [Homo sapiens] ABC (ARA001LB (ARA001LB [Homo sapiens] ABC (ARA001LB (ARA001LB (ARA001LB (ARA001LB (ARA001LB (ARA00		79845694 ((1983, 198	 Novel Protein sim. GBank gi[2105049]emb CAB08835] - (Z95436) hypothetical protein Rv3645 [Mycobacterium luberculosis] 	Contains protein domain (PF00211) - Adenylate and Guanylate cyclase catalytic domain	UNCLASSIFIED	264508, 264593
11090390 (1987, 1988) Novel Protein sim. GBank gi]3329297 (AE001355) - Reductase Reductase, Large Chain [Chlamydia Ribonucleoside Reductase, Large Chain [Chlamydia Submucleoside Reductase, Large Chain [Chlamydia Submucleoside Reductase, Large Chain (AF143946) SNF2 and others N-terminal domain (AF00176) - helicase gi 5106572[gb AaD39760.1 AF14394 - (AF143946) SNF2 and others N-terminal domain (AF00176) - helicase gi 5106572[gb AaD39760.1 AF14394 - (AF143946) SNF2 and others N-terminal domain (AF00156) - helicase gi 5106572[gb AaD39760.1 AF17010427] - Contains protein domain (PF00056) - transport transporter ATP-binding protein [Archaeoglobus fulgidus] ABC transporter ATP-binding protein sim. GBank gi 23530[sp P04929]HRPX_PLALO - HISTIDINE-RICH GLYCOPROTEIN PRECURSOR GLYCOP	Γ	10814053 (1985, 198	+-			264907
94321911 (1989, 1990) Novel Protein sim. GBank gil2649101 (AE001001) - ABC (1991, 1992) Novel Protein sim. GBank gil2649101 (AE001001) - ABC (Contains protein domain (PF00005) - transporter, ATP-binding protein fachaeoglobus fulgidus] ABC transporter GLYCOPROTEIN PRECURSOR		11090590 ((1987, 198	8) Novel Protein sim. GBank gij3329297 (AE001355) - Ribonucleoside Reductase, Large Chain (Chlamydia trachomatic)		reductase	264602
### 1397390 (1995, 1996) Novel Protein sim. GBank gil2911PT9 (AC004227) - ###################################	T	1 101010	1000	Manual Design of a Cook	Contains profein demain (PE00176)	helicase	18108398 65274572 22278996 264490.
91013745 (1991, 1992) Novel Protein sim. GBank gi[2811719 (AC004227) - Contains protein domain (Proteins sim. GBank gi[2649101 (AE001001) - ABC Contains protein domain (Proteins sim. GBank gi[2649101 (AE001001) - ABC Contains protein domain (Proteins sim. GBank gi[2649101 (AE001001) - ABC Contains protein domain (Protein sim. GBank gi[2649101 (AE001001) - ABC transporter gi[123530]sp[P04929]HRPX_PLALO - HISTIDINE-RICH GLYCOPROTEIN PRECURSOR		9432101	66	두윤 [SNF2 and others N-terminal domain		264905, 29331827, 28146498, 264508, 264909, 29331827, 28146498, 264508, 264908, 264509, 264591, 264692, 60432229, 60431735, 33657402, 264595, 264758, 21906754, 265010, 265017, 265018, 264605, 264760, 26448, 264763, 264768, 21906765, 21906769, 55811957, 264692, 264693, 264629, 3569643, 55811576, 35695855, 264558, 284556, 264558, 284553, 284563, 284630, 2846
80503347 (1993, 1994) Novel Protein sim. GBank gil2649101 (AE001001) - ABC Contains protein domain (PF00005) - transport transporter transporter, ATP-binding protein [Archaeoglobus fulgidus] ABC transporter ABC transporter (1397390 (1995, 1996) Novel Protein sim. GBank gil123530lsp P04929 HRPX_PLALO - HISTIDINE-RICH GLYCOPROTEIN PRECURSOR		91013745 ((1991, 199	2) Novel Protein sim. GBank gi 2911719 (AC004227) - KIA001LB [Homo sapiens]	Contains protein domain (PF00595) - PDZ domain (Also known as DHR or GLGF).	pnospnatase	83373044, 23331824, 283378, 33611337, 56526486, 87168518, 264910, 264906, 264565, 264566, 264693, 264766
11397390 (1995, 1996) Novel Protein sim. GBank gij123530[sp P04929]HRPX_PLALO - HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	,	80503347	(1993, 199	Novel Protein sim. GBank gi[2649101 (AE001001) - ABC transporter, ATP-binding protein [Archaeoglobus fulgidus]	Contains protein domain (PF00005) - ABC transporter	transport	35696286, 22278997, 22278999, 264508, 264905, 264908, 265010, 264600, 264602, 264605, 264688, 264769, 265021, 264565, 18108391
	1]	11397390	(1995, 199	×' <u>&</u>			264595

							itase 35696052, 264508, 265008, 265009, 264769, 18108387, 264563	associated 264593	UNCLASSIFIED 52646842, 33657402, 33657023, 18108379, 55811576, 264631, 264556, 264557, 264559, 18108388, 264566		-	UNCLASSIFIED 265017
-00040) - struct	esterase	F00496) - transport e-binding		UNCLASSIFIED		synthase	F00459) - phospha family	F00122) - ATPase	UNCLAS	transport	helicase	UNCLA
Contains protein domain (PF00040) - struct Fibronectin type II domain		Contains protein domain (PF00496) - transport Bacterial extracellular solute-binding proteins, family 5					Contains protein domain (PF00459) - phosphatase Inositol monophosphatase family	Contains protein domain (PF00122) - ATPase_associated 264593 E1-E2 ATPase				
95418879 (2029, 2030) Novel Protein sim. GBank gil4159995 (AF063095) - SELIL [Mus musculus]	79559694 (2031, 2032) Novel Protein sim. GBank gi 2506969 sp P41407 ACPD_ECOLI - ACYL CARRIER PROTEIN PHOSPHODIESTERASE (ACP PHOSPHODIESTERASE)	Novel Protein sim. GBank gi 5103943 dbj BAA79259.1 - (AP000059) 802aa long hypothetical oligopeptide-binding protein oppA [Aeropyrum pernix]	80072430 (2035, 2036) Novel Protein sim. GBank gil4493973 emb CAB39032.1 - (AL034559) predicted using hexExon; MAL3P7.14 (PFC0925w), Hypothetical protein, Ien: 489 aa [Plasmodium falciparum]			37036243 (2041, 2042) Novel Protein sim. GBank gil4633807[gb]AAD26859.1[AF12779 - (AF127795) trehalose biosynthetic enzyme TreY (Rhizobium leguminosanum by. viclael	80502627 (2043, 2044) Novel Protein sim. GBank gi 1781230 emb CAB06277 - (Z83867) hypothetical protein Rv3137 [Mycobacterium tuberculosis]	11399341 (2045, 2046) Novel Protein sim. GBank gil3777495 (U92083) - calcium Irransporting ATPase [Pichia angusta]		79644200 (2049, 2050) Novel Protein sim. GBank gij3483045 emb CAA20556 - (AL031371) putative transport system permease protein IStreptomyces coelicolori	80025946 (2051, 2052) Novel Protein sim. GBank gil1174922[sp]Q02322[UVRD HAEIN - DNA HELICASE II	17659234 (2053, 2054) Novel Protein sim. GBank gil4757728IrefINP_004886.1lpAGTA - anoidensin/vasopressin receptor All/AVP-like
5418879 (2029, 2030)	9559694 (2031, 2032)	1069213 (2033, 2034)	0072430 (2035, 2036)	11703607 (2037, 2038)	1020 80234432 (2039, 2040)	37036243 (2041, 2042)	30502627 (2043, 2044)	1399341 (2045, 2046)	80057129 (2047, 2048)	79644200 (2049, 2050)	30025946 (2051, 2052)	17659234 (2053, 2054
1015	1016 7	1017	1018	1019	1020	1021	1022	1023 1	1024	1025	1026	1027

300,					
970	(0502, 2050)	1029 / 2029/926 (2035, 2030) Novel Protein sim. GBark gijz / 91409[emb](JAA 6003] - (AL021184) acn [Mycobacterium tuberculosis]	Contains protein domain (PF00330) - UNCLASSIFIED Aconitase family (aconitate	UNCLASSIFIED	264600
3			hydratase)		
870	_			UNCLASSIFIED	264595
9	88095343 (2059, 2060)			UNCLASSIFIED	264907, 264908, 264510, 264512, 265009, 265010, 265011, 264600, 264602, 264603,
1031	95289117 (2061, 2062)			UNCLASSIFIED	264905, 264709, 16106372, 16106374 264905, 264906, 264909, 264595, 264692,
900					264630, 264634, 264638
1032	94673275 (2063, 2064)	946 / 32 / 5 (2063, 2064) Novel Protein sim. GBank gi 4503895 ref NP_000145.1 pGALK - galactokinase 1		UNCLASSIFIED	264689
1033	86464818 (2065, 2066)	86464818 (2065, 2066) Novel Protein sim. GBank gil2982990 (AE000682) -		UNCLASSIFIED	35696052, 264906, 264510, 18108354.
		hypothetical protein [Aquifex aeolicus]			264687, 264769, 264689, 60431602, 18108385, 264486
1034	79245937 (2067, 2068)	79245937 (2067, 2068) Novel Protein sim. GBank gil405895 (U00007) - methionyl-		UNCLASSIFIED	264906
4025	7006 0306 73663	many symmetrical promising confi			
				UNCLASSIFIED	264692
_	_				264905, 66712502, 264908, 264766
1037	87896058 (2073, 2074)			UNCLASSIFIED	29331824, 264909, 60433438, 265019
	20481015 (2075, 2076)	Novel Protein sim. GBank gi 790819 (L39891) - polycystic kitnev dispessassociated protein Homo seriene	Contains protein domain (PF01477) -		264604, 264634
4030	87360034 (3077 3078)	97260034 (2077 2070) Maint Datain aim OB -1 -10006007 (AF0000007)	ויביוג מסוומוו		
	(2017, 2010)	inovel Protein Sim. GBank gitzbubbo/ (AFubuuzz) - 24 Equine herpesvirus 4		UNCLASSIFIED	264092, 264093, 264094, 264683, 264689, 263967
1040	80026840 (2079, 2080)	80026840 (2079 2080) Novel Protein sim GBank nij2352095 (192022) - DNA	Contains protein domain (DE01131) isomeraso	icomoraco	100002
		[topolsomerase [Fervidobacterium islandicum]	Prokaryotic DNA topoisomerase	2	
1041	10156682 (2081, 2082)	10156682 (2081, 2082) Novel Protein sim. GBank gil3256535 db BAA29218.1 -		kinase	264907
		(AP000001) 301aa long hypothetical 2-phosphoglycerate			
1		Airiasa Ir yiococcus ilorikosiilij			
1042	11084375 (2083, 2084)	11084375 (2083, 2084) Novel Protein sim. GBank gi 2058299 emb CAA66953 - (X98309) ARI protein [Drosophila melanogaster]			264605
1043		80057136 (2085, 2086) Novel Protein sim. GBank gil1870167 emb CAA70125 -	Contains protein domain (PF00005) - transport	transport	264565, 264567
		וו מספרול ווופוע לפוובלומוול מפין בחבמוו	ADC transporter		
5		57445403 (2007, 2009)		UNCLASSIFIED	265006, 264602, 265017
		Novel Protein sim. Grank gijobššišujembjCAB52053.1 -			29331825, 264637
1046	_	11754862 (2091, 2092) Novel Protein sim. GBank gil854065jembjCAA58337j		helicase	264686
_		(A63413) U86 Human nerpesvirus 6			
	3/036258 (2093, 2094)	3/035258 (2093, 2094) Novet Protein sim. GBank gil4210471 dbj BAA74535.1 - (AB019033) orfSA [Pseudomonas sp.]		UNCLASSIFIED	264769
1048	79186400 (2095, 2096)	Novel Protein sim. GBank gil3413419 emb CAA20279 -	Contains protein domain (PF00220) - UNCLASSIFIED	UNCLASSIFIED	264687
		(AL031232) hypothetical protein SC10H5.07 [Streptomyces			
		coenicolor	terminal Domain		
3	01/33100 (2037, 2038)	01/33100 (2097, 2098) Nover Protein Sim. GBank oil5051636lob AAD38326 1 AED7372 - (AED73727 FH		UNCLASSIFIED	264905, 264634
		domain-binding mitotic phosphoprotein [Homo sapiens]			
1050	79471521 (2099, 2100)			UNCLASSIFIED	264686

1051	1051 [80475471 (2101, 2102)			UNCLASSIFIED	18108374 264769 265010 265011 264601
					265009, 264604, 264605, 264636, 18108351, 264692
1052		82442962 (2103, 2104) Novel Protein sim. GBank	Contains protein domain (PF00389) - dehydrogenase	dehydrogenase	264508, 264762, 264687, 264486
		gi[3123275 sp P35136 SERA_BACSU - D-3-	D-isomer specific 2-hydroxyacid		
	_	PHOSPHOGLYCERATE DEHYDROGENASE (PGDH)	dehydrogenases		
1053		94851640 (2105, 2106) Novel Protein sim. GBank gij5441319 emb CAB46717.1 -			264686, 18108374, 29331824, 83373044,
		(AL034396) dJ1158B12.1 (zinc finger, X-linked, duplicated			21906754, 52645156, 56182435, 264689,
		A) [Homo sapiens]			29331827, 27486261, 35696052, 21908765, 1
					35696423, 21906/68, 561625/5, 21906/69,
				٠	55811957, 87168518, 35696286, 22278997,
_					265020, 265011, 265021, 265022, 265007.
					265018, 22279000, 22279002, 264482,
				-	264906, 52644150, 264909, 264288,
					29331822, 52645080, 264766
1054				UNCLASSIFIED	264686
1055		80594138 (2109, 2110) Novel Protein sim. GBank	Contains protein domain (PF00270) - helicase	helicase	264907, 264602, 264681, 264288, 21906768,
		gi[5052508 gb AAD38584.1 AF14560 - (AF145609) BeDNA GH02833 [Dresnahir melanometer]	DEAD/DEAH box helicase		33657109, 55810764, 35695855, 264631
1058	_	17882319 (2111 2112) Invole Protein sim GRank dit 1021676 Intellige A 251581		rnapolymerace	264906
}	_	(Coecas) DNA actions given 20 factor (Decidence)		napoly include	
		(Locossy) KNA polymerase sigma-70 factor (r seudomonas fluorescens)			
1067	85567216 (2113 2114)	85557945 (2412 21144) Mound Deutsin sim Chark siltanens (150209) Mo		CHELEN	000000
3		definition line found [Caenorhabditis elegans]		ORCEASSIFIED	700,607
1058	80376576 (2115, 2116)	-			264764
1059		94662754 (2117, 2118) Novel Protein sim. GBank		transcriptfactor	35696052, 35695855, 265009, 264636
		MOITGOOT STANKE MAN TO TO ANY CONTROL ON			
		ELONGATION FACTOR GREA (TRANSCRIPT CLEAVAGE			
		FACTOR GREA)	:		
1060		79481169 (2119, 2120) Novel Protein sim. GBank		glycoprotein	29146499, 264681, 264683, 264687
		gi[2499087]sp[Q09332]UGGG_DROME - UDP-			
		GLUCOSE: GLYCOPROTEIN GLUCOSYLTRANSFERASE			
		PRECURSOR (DUGT)			
1061	11034025 (2121, 2122)	11034025 (2121, 2122) Novel Protein sim. GBank gij90254 pir A28334 - protein-		phosphatase	264634
		[tyrosine-phosphatase (EC 3.1.3.48) Ly-5 precursor (B-cell	,		
	_	variant) - mouse			
1062		39567937 (2123, 2124) Novel Protein sim. GBank		dehydrogenase	264593
		gi 3334200 sp O49954 GCSP_SOLTU - GLYCINE			
		DEHYDROGENASE (DECARBOXYLATING) PRECURSOR			
		(GLYCINE DECARBOXYLASE) (GLYCINE CLEAVAGE			
		SYSTEM P-PROTEIN)			
1063	8490481 (2125, 2126)	Novel Protein sim. GBank			264508
		gi[2499966 sp Q41228 PSE1_NICSY - PHOTOSYSTEM			
		REACTION CENTRE SUBUNITIVA PRECURSUR (PSI-ELA)			
1064	78891783 (2127, 2128)	78891783 (2127, 2128) Novel Protein sim GBank pil82654 lpiril IA0086 - 10K zein			265007 265008 18108351 18108385
		precursor - maize			
			1		

1065	80021208 (2129, 2130)	1065 80021208 (2129, 2130) Novel Protein sim. GBank gil212098 pir S70682 -		transferase	264600, 264602, 264689
1066	17896879 (2131, 2132)	17896879 (2131, 2132) Novel Profession GBank gli2506362[sp[P15042]DNL_J_ECOLI - DNA LIGASE (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (NAD+))		synthase	265009
1067	10132178 (2133, 2134)				264909
1068		82062057 (2135, 2136) Novel Protein sim. GBank gil4007669 emb CAA22355 - (AL034443) putative oxidoreductase [Streptomyces coelicolor]	Contains protein domain (PF00248) - reductase Aldo/keto reductase family		264688, 18108362, 264558, 264600, 264760
1069		83002954 (2137, 2138) Novel Protein sim. GBank gil4589484 dbj BAA76770.1 - (AB023143) KIAA0926 protein [Homo sapiens]		UNCLASSIFIED	264604
1070				UNCLASSIFIED	264604, 264760
		gij120304[sp[P15932]FLGK_SALTY - FLAGELLAR HOOK- ASSOCIATED PROTEIN 1 (HAP1)			
107		20710589 (2141, 2142) Novel Protein sim. GBank gil1750397 (U81261) - glutamate synthase large subunit [Pseudomonas aeruginosa]			264602
1072	82356540 (2143, 2144)			UNCLASSIFIED	264687, 264688, 21906764, 35696052,
					35895917, 35895855, 264600, 264601. 264602, 265009, 264605, 264508, 264905, 264690, 264906, 264762, 264628, 264766
1073	79814400 (2145, 2146)				264909
1074	_	80105992 (2147, 2148) Novel Protein sim. GBank gil477532 pir A49175 - Motch B protein - mouse (fragment)	Contains protein domain (PF00008) - synthase EGF-like domain	synthase	264906
1075		81850293 (2149, 2150) Novel Protein sim. GBank qil3893109lemb CAA76940 -		UNCLASSIFIED	56994075, 22278998, 264594, 264757,
		(Y17920) CALO protein [Drosophila melanogaster]			264596, 265018, 265019, 264681, 264369. 284688, 285020, 18108364, 18108374
1076		80477264 (2151, 2152) Novel Protein sim. GBank		ATPase_associated	264769
		gi[1176203[sp[P4642]YHCM_ECOLI - HYPOTHETICAL 43.1 KD PROTEIN IN RPLM-HHOA INTERGENIC REGION (F375)			
1077		79831334 (2153, 2154) Novel Protein sim. GBank gil4033487 sp Q44472 TUD4_AGRVI - PUTATIVE HYDROXYPYRUVATE REDUCTASE			264905
1078	20288874 (2155, 2156)				264600
1079		80494518 (2157, 2158) Novel Protein sim. GBank gil3413828 emb CAA20296 - (AL031260) hypothetical protein SC9A10.09 [Streptomyces coelicolor]		UNCLASSIFIED	18108394, 264769, 264634, 264636
1080	11767188 (2159, 2160)	-		UNCLASSIFIED	264684
1081	94747080 (2161, 2162)				83373044, 265019, 22279002, 264482, 18108351, 264682, 264908, 264693, 264487
1082	81490656 (2163, 2164)			UNCLASSIFIED	264758, 264768, 264769, 21906767, 264511,
					264910, 264634, 264635, 264905, 264955, 264906, 264637, 264907, 264908, 264764,
					264638, 20281099, 264700, 264393

83	87446717 (2165, 2166)	1083 87446717 (2165, 2166) Novel Protein sim. GBank gil 1722945 sp Q10523 Y01N_MYCTU - HYPOTHETICAL		UNCLASSIFIED	60424179, 264905, 264906, 264510, 60432229, 264759, 87168474, 264605,	
		44.6 KD PROTEIN CY427.23			264769, 264689, 18108364, 18108376, 35695855, 264636	
1084	37799306 (2167, 2168)	37799306 (2167, 2168) Novel Protein sim. GBank gil418384 sp P32057 WCAL_ECOLI - PUTATIVE COLANIC ACID BIOSYNTHESIS GLYCOSYL TRANSFERASE WCAI		UNCLASSIFIED	264769	
1085	86475368 (2169, 2170)	heat	Contains protein domain (PF00118) - eph TCP-1/cpn60 chaperonin family	eph	60432229, 264687	
_	79608269 (2171, 2172)		Contains protein domain (PF00238) - ribosomalprot Ribosomal protein L14		264486	
	79603979 (2173, 2174)	31 - protein)	Contains protein domain (PF00385) - helicase 'chromo' (CHRromatin Organization MOdifler) domain	helicase	29331827, 264693	
	79854963 (2175, 2176)	79854963 (2175, 2176) Novel Protein sim. GBank gi 2983155 (AE000693) - phosphoglucomulase/phosphomannomulase [Aquifex aeolicus]			264905, 264601, 18108387	
1089	80216800 (2177, 2178)	80216800 (2177, 2178) Novel Protein sim. GBank gil4981768 gb AAD36290.1 AE00177 - (AE001778) NADH dehydrogenase, 30 kDa subunit, putative [Thermotoga maritima]	Contains protein domain (PF00329) - UNCLASSIFIED Respiratory-chain NADH dehydrogenase, 30 Kd subunit		264488, 264511, 265011, 264682, 264768, 264689, 21906764, 35695917, 265020, 32833986, 18108370, 35695855	
1090	11083825 (2179, 2180)	11083625 (2179, 2180) Novel Protein sim. GBank gil4007680 emb CAA22366 - (AL034443) putative oxidoreductase [Streptomyces coelicolor]			264604	
1091	12917471 (2181, 2182)	12917471 (2181, 2182) Novel Protein sim. GBank gi[2495562[sp P77239]YLCD_ECOLI - HYPOTHETICAL 44.3 KD PROTEIN IN NFRB-PHEP INTERGENIC REGION PRECURSOR		UNCLASSIFIED	264637	
	80252286 (2183, 2184)	Г			264566	
1093	80496304 (2185, 2186)	-	Contains protein domain (PF00730) - nuclease Endonuclease III	nuclease	264769, 35695917, 35695855, 264600, 264602, 264603, 264605, 18108351	
1094	10880972 (2187, 2188)	10880972 (2187, 2188) Novel Protein sim. GBank gil1001642 dbj BAA10373 - (D64002) dGTP triphosphohydrolase [Synechocystis sp.]		UNCLASSIFIED	264686	
		Novel Protein sim. GBank gil4585587[emb CAB40855.1] - (AL049628) putative adenine glycosylase [Streptomyces coelicolor]	Contains protein domain (PF00455) - Inuclease Bacterial regulatory proteins, deoR family		264906, 265007, 264595, 264600, 264602, 264603, 264604, 264605, 264762, 264766, 264769, 264769, 264482, 18108387, 60432113, 264482, 264486	
		80025977 (2191, 2192) Novel Protein sim. GBank gij115001jspjP19206jBIOB_BACSH - BIOTIN SYNTHASE (BIOTIN SYNTHETASE)		synthase	264600, 264602, 264603, 264604, 264605, 35695917, 264692, 264631	
1097	79239560 (2193, 2194)			UNCLASSIFIED	265019	
	79186424 (2195, 2196)	79186424 (2195, 2196) Novel Protein sim. GBank gij114135[sp]P08205JARGA_ECOLI - AMINO-ACID ACETYLTRANSFERASE (N-ACETYLGLUTAMATE SYNTHASE) (AGS)		synthase	264687	

	264259, 264636	264769		264757	D 29331822, 21906754, 264555, 284556, 264558, 22279002	264566	Γ	264908, 264769	264907, 264510, 264511, 264600, 264602, 264605, 264768, 264769	264602, 264605, 264636	264908, 264593, 265010, 264601, 264603, 264604, 264605, 264682, 264769, 264693, 264636	18108370, 264557	264908		a 264636	l	264908
UNCLASSIFIED	desaturase	isomerase	UNCLASSIFIED	struct	UNCLASSIFIED		UNCLASSIFIED		transport	helicase				UNCLASSIFIED	dehydrogenase	UNCLASSIFIED	
		Contains protein domain (PF01396) - isomerase Topoisomerase DNA binding C4 zinc finger															
1099 39523838 (2197, 2198) Novel Protein sim. GBank gij3915144 sp 033017 TRMD_MYCLE - TRNA (GUANINE- N1)-METHYLTRANSFERASE (M1G- METHYLTRANSFERASE) (TRNA [GM37] METHYLTRANSFERASE)	85736571 (2189, 2200) Novel Protein sim. GBank gij3023255 sp Q64420 ACOD_MESAU - ACYL-COA DESATURASE (STEAROYL-COA DESATURASE) (FATTY ACID DESATURASE) (DELTA(9)-DESATURASE)	80491857 (2201, 2202) Novel Protein sim. GBank gi 1174735 sp P43012 TOP1_HAEIN - DNA TOPOISOMERASE I (OMEGA-PROTEIN) (RELAXING ENZYME) (UNTWISTING ENZYME) (SWIVELASE)			95003115 (2207, 2208) Novel Protein sim. GBank gil2935448 (AF048976) - synaptic ras GTPase-activating protein p135 SynGAP (Rattus norvegious)				80440616 (2215, 2216) Novel Protein sim. GBank gil1173421 sp P43416 SECY_STRSC - PREPROTEIN TRANSLOCASE SECY SUBUNIT	80064615 (2217, 2218) Novel Protein sim. GBank gi 2995310 emb CAA18338 - (AL022268) putative ATP-dependent helicase Streptomyces coelicolor		80071744 (2221, 2222) Novel Protein sim. GBank gijz622039 (AE000868) - type I restriction modification system, subunit S [Methanobacterium thermoautotrophicum]		82456352 (2225, 2226) Novel Protein sim. GBank gij3218376jembjCAA19628j - (AL023862) putative oxidoreductase [Streptomyces coelicolor]	14998014 (2227, 2228) Novel Protein sim. GBank gi 1083428 pir S54876 - NAD(P)+ transhydrogenase (B-specific) (EC 1.6.1.1) precursor - mouse		
39523838 (2197, 2198) F	85736571 (2199, 2200) 1	80491857 (2201, 2202) 1 9 T	79777614 (2203, 2204)	81897259 (2205, 2206) 	95003115 (2207, 2208) s s	80255121 (2209, 2210)	79314110 (2211, 2212)	80470019 (2213, 2214)	80440616 (2215, 2216) h	80064615 (2217, 2218) (4 (1)	80503554 (2219, 2220)	80071744 (2221, 2222)	95010088 (2223, 2224)	82456352 (2225, 2226) h	14998014 (2227, 2228) N N P	11765583 (2229, 2230)	1116 79841152 (2231, 2232)
1099			1102	1103	1104	1105	1108			1109	1110			1113	1114	1115	1116

1119 195303465 (2233, 2234) Nove Protein stim. Glank gji255955 emb CA64089] - Contains protein domain (PF00270) - helicase	18108392, 264488, 263994, 264489, 18108398, 56182575, 22278996, 35696286, 56994075, 22278995, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 264259, 29331822, 56182181, 29331824, 66714117, 29331825, 29331826, 264309, 2644004, 56182435, 264500, 2644004, 56182435, 264500, 264501, 264599, 264590, 264500, 264501, 264599, 264590, 264601, 264602, 26401, 264601, 264602, 26401, 264602, 26401, 264060, 264601, 264602, 264603, 264017, 264061, 264602, 264603,	264691	264907	29331827, 265018, 265019, 264681, 265021, 60170615, 18108387	264683, 264632, 18108388	264639, 264563	264693, 27486265	264909	264910	264905, 264909, 264910	264682	264511
95305465 (2233, 2234) Novel Protein sim. GBank gij3255965jembj(CA694089j - (Z70200) U5 snRNP-specific 200kD protein [Homo sapiens] 79642463 (2236, 2236) 79642463 (2239, 2236) 79471716 (2241, 2242) Novel Protein sim. GBank gij5420387lembjCA846579.1-(Genochabdisis elegans] 79671716 (2242, 2242) Novel Protein sim. GBank gij64450 (U67864) · MEX-3 (Genochabdisis elegans] 79637119 (2245, 2246) Novel Protein sim. GBank gij98800jprij[S17768 - 3-dehydroquinate synthase (EC 4.6.1.3) · Mycobacterium taberculosis 79811596 (2249, 2250) Novel Protein sim. GBank gij9800jprij[S17768 - 3-dehydroquinate synthase (EC 4.6.1.3) · Mycobacterium taberculosis 79811596 (2249, 2250) Novel Protein sim. GBank gij98002704jembj(CA844358.1)-(AL242630) Novel Protein sim. GBank gij5002704jembj(CA844358.1)-(AL242630) DNA polymerase i [Methylopacterium sp. DNA)	helicase	UNC! ASSIFIED	UNCLASSIFIED	collagen	UNCLASSIFIED	UNCLASSIFIED	synthase	UNCLASSIFIED	UNCLASSIFIED	eph	UNCLASSIFIED	polymerase
95305465 (2233, 2234) Novel Protein sim. GBank gi (Z70200) U5 snRNP-specific 79563326 (2236, 2236) 795642463 (2239, 2240) Novel Protein sim. GBank gi (A-1243459) proteophosphog 79471716 (2241, 2242) Novel Protein sim. GBank gi (Caenorhabditis elegans) 79456246 (2243, 2244) 79557861 (2243, 22245) Novel Protein sim. GBank gi dehydroquinate synthase (Einberculosis 79757861 (2242, 2256) Novel Protein sim. GBank gi 79758914 (2251, 2252) Novel Protein sim. GBank gi 7975885 (2253, 2254) Novel Protein sim. GBank gi 78384885 (2255, 2256) Novel Protein sim. GBank gi (A-1242630) DNA polymerase	Contains protein domain (PF00270) - DEAD/DEAH box helicase				Contains protein domain (PF00013) - KH domain							Contains protein domain (PF00476) - DNA potymerase family A
	Novel Protein sim. GBank gij3255965 emb CAA94089j - (Z70200) U5 snRNP-specific 200kD protein [Homo sapiens]			Novel Protein sim. GBank gi 5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]	Novel Protein sim. GBank gi 1644450 (U67864) - MEX-3 [Caenorhabditis elegans]		Novel Protein sim. GBank gil98800 pir S17768 - 3- dehydroquinate synthase (EC 4.6.1.3) - Mycobacterium tuberculosis			7,6		5002704(emb CAB44358.1] - 9 I [Methylobacterium sp. DM4]
10010010 to 101m total of the l	95305465 (2233, 2234)	_				79456246 (2243, 2244)	79637119 (2245, 2246)	79811596 (2247, 2248)	79757861 (2249, 2250)	79758914 (2251, 2252)	И	

ED 265011, 264766	Т	ED 14606286 22278008 20331828 264603			264604, 264605, 264760, 32833986, 18108374		ED 264512, 264534	264508, 264600, 264602, 264603, 18108376	ED 264906, 264908	22278998, 22278999, 35696052, 264907, 265009, 60433356, 264596, 265010, 264448, 264682, 264767, 264689, 265020, 264692, 55811576, 35695855, 264631, 264632, 22279002	264908	ED 264693					ED 264635, 264636, 264907, 264593, 264908, 264566, 264909	ED 264112	ED 264769, 264689, 35696286, 264760, 264905, 264486, 264559	264557	284591	264591
UNCLASSIFI	JINO ACOIC	INCI ASSIETED		UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	collagen	struct	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIF	oxidase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		kinase	
Contains protein domain (PF00170) - UNCLASSIFIED	ozir transcription factor									Contains protein domain (PF00515) - collagen TPR Domain						Contains protein domain (PF00090) - oxidase Thrombospondin type 1 domain						
80422480 (2257, 2258) Novel Protein sim. GBank gil5689485 dbj BAA83026.1 -	(Abuzoss) NAMIU 4 protein [Homo sapiens]	80055391 (226) 2200) Novel Protein cim CRank	gil4981328lgblAAD35881.1JAE00174 - (AE001747) bioY protein [Thermotoga maritima]	82062248 (2263, 2264) Novel Protein sim. GBank gil 1841552 (U89336) - unknown	[Homo sapiens]			Novel Protein sim. GBank gil4539171 emb[CAB39700.1] - (AL049485) conserved hypothetical protein [Streptomyces coelicolor]	79842052 (2271, 2272) Novel Protein sim. GBank gil4982454lgblAAD36931.1JAE00182 - (AE001823) ATP- dependent protease LA, putative [Thermotoga maritima]	90931557 (2273, 2274) Novel Protein sim. GBank gil4972746[gbJAAD34768.1] - (AF132180) unknown [Drosophila melanogaster]	79841163 (2275, 2276) Novel Protein sim. GBank gil731607 sp P38739 YHC8_YEAST - HYPOTHETICAL 63.8 KD PROTEIN IN GUT1-RIM1 INTERGENIC REGION PRECURSOR	78633561 (2277, 2278) Novel Protein sim. GBank gij3650031 (AC005396) - putative protine-rich cell wall protein [Arabidopsis thaliana]				87762158 (2285, 2286) Novel Protein sim. GBank gi]3928000 emb CAA05880] - (AJ003125) procollagen I N-proteinase [Homo sapiens]	_				Novel Protein sim. GBank gi[2499003 sp P76422 THID_ECOLI - PHOSPHOMETHYLPYRIMIDINE KINASE (HMP- PHOSPHATE KINASE) (HMP-P KINASE)	1149 11607438 (2297, 2298) Novel Protein sim. GBank gil2896734 emb CAA17213.1 - (AL021897) hypothetical protein Rv1097c [Mycobacterium
80422480 (2257, 2258)	70420151 (2250 2250)	RDD55391 (2261 2262)		82062248 (2263, 2264)		17290437 (2265, 2266)	80235376 (2267, 2268)	80029393 (2269, 2270)	79842052 (2271, 2272)	90931557 (2273, 2274)	79841163 (2275, 2276)	79633561 (2277, 2278)	39480358 (2279, 2280)	79638019 (2281, 2282)	19635848 (2283, 2284)	87762158 (2285, 2286)	80088988 (2287, 2288)	14610262 (2289, 2290)	82062092 (2291, 2292)	80071761 (2293, 2294)		11607438 (2297, 2298)
1129				1132		1133	1134	1135	1136	1137	1138	1139	Т		1142	1143	1144	1145	1146	1147		1149

ELICATION CARE CALL SANCE	74 (2299, 2300	Novel Protein sim. GBank gij2895095 (AF011337) - putative	IA1	Pase_associated	ATPase_associated 264488, 35696286, 264907, 264908, 264909,
24655 gb AAD16978 -		E1-E2 ATPase [Mus musculus]			264910, 264593, 264596, 264758, 264764.
137275 emb CAB0731.1 - polymerase 2 13					264766, 264768, 264693, 264628, 60431850,
4324653[gbtAAD16978] - 18 lapha subunit 18674275[emb]CAB07311.1 - 18747[PS.32467]; CDN EST 18 gene; CDNA EST yk404c10.5 18 gene; CDNA EST yk404c10.5 19 Gondains protein domain (PF00122) - transport 19 ECOLI - ZINC - 19 ECOLI - HYPOTHETICAL 34.0 19 ECOLI - HYPOTHETICAL 34.0 19 ECOLI - HYPOTHETICAL 34.0 19 EA3342[dbj BAA27380] - 19 EA3342[dbj BAA27380] - 19 EA3342[dbj BAA27380] - 19 EA4563 - (AF145632) - 19 EA4563 - (AF145632) - 19 EA3342[dbj BAA76766.1] - 19 EA3345[dbj BAA76766.1] - 19 EA3345[dbj BAA76766.1] - 19 EA3945[dbj BAA76766.1] - 10 EA3945[dbj BAA767676.1] - 10 EA3945[dbj BAA767676.1] - 10 EA3945[dbj BAA767676.1] - 10 EA3945[dbj BAA767676.1] - 10 EA3945[dbj BA					264564, 264566, 264567
### ### ### ### ### ### ### ### ### ##	302		od		264595
gij3874275[emb]CAB07311.1 - SaneInder; Similarity to Yeast low SaneInder; Similarity to Yeast low Ashering to Yeast low a this gene; cDNA EST yk404c10.5 ### (PS:32467); CDNA EST yk404c10.5 is gene; cDNA EST yk404c10.5 UNCLASSIFIED ONCLASSIFIED ONCLA		(AFTUOL91) DIVA polymerase in alpha subumit (Streptomyces coelicolor)			
Hart PS:32467; cDNA EST Low Hart PS:3247; contains protein domain (PF00122) Lansport Hart PS:3247; contains Low Hart PS:3247; contains Low Hart PS:3267; contains Low Hart Hart Low Hart PS:3267; contains Low Hart Hart Low Hart PS:3267; contains Low Hart Hart Low Hart Hart Low Hart Low Hart Hart Low Hart Hart Hart Hart Hart Hart Hart Hart Hart Hart Hart	8	Novel Protein slm. GBank gij3874275[emb[CAB07311.1] -	(16)		264488, 22278998, 264905, 264629, 264486
In this gene; cDNA EST yk404c10.5 Its gila240315[bib]BAA74936.1] - Contains protein domain (PF00122) - transport E (ZN(II)-TRANSLOCATING P- E-E		(292825) predicted using Genefinder; Similarity to Yeast low		•	
is gene; cDNA EST yk404c10.5 gig4240315[dbj BAA74936.1] - Contains protein domain (PF00122) - transport E (ZN(II)-TRANSLOCATING P- E1-E2 ATPase		FMBI : C12555 comes from this gene: cDNA EST			
### 191240315 dbj BAA74936.1 - JuncLASSIFIED UNCLASSIFIED Lontains protein domain (PF00122) - transport E (ZN(II)-TRANSLOCATING P - E1-E2 ATPase UNCLASSIFIED LECOLI - ZINC E1-E2 ATPase UNCLASSIFIED LONCLASSIFIED LONCLASSIFI		yk404c10.3 comes from this gene; cDNA EST yk404c10.5		THE STATE OF THE S	
Michael Sapiens Contains protein domain (PF00122) - Iransport		comes from thi			
9ij4240315jdbjjBAA74936.11- 2te (ZN(II)-TRANSLOCATING P- E (ZN(II)-TRANSLIFED P- INCLASSIFIED P- E (ZN(II)-TRANSLOCATING P- E (ZN(II)-TRANSLIFED P- INCLASSIFIED P- E (ZN(II)-TRANSLIFED P- INCLASSIFIED P- E (ZN(II)-TRANSLIFED P- INCLASSIFIED P- E (ZN(II)-TRANSLOCATING P- INCLASSIFIED P- INCLASSI	38			I	264910, 264555, 264557
UNCLASSIFIED	2308		5		265008
Contains protein domain (PF00122) - transport	310		Ō	SIFIED	264690, 264636
N_ECOLI - ZINC	2312		ains protein domain (PF00122) - tra	ansport	264603
Coccasion Cocc		N_ECOLI - ZING- E (ZN(II)-TRANSLOCATING P.	2 ATPase		
2 ECOLI - HYPOTHETICAL 34.0 3N INTERGENIC REGION 9S_MYCTU - HYPOTHETICAL 27C 27C 27C 27C 27C 27C 27C 27	314	Novel Protein sim, GBank	5	NCLASSIFIED	264906, 264907, 264758, 264766, 264769,
Samuel Marca Mar		고Ж			264689, 264638, 264566
SamyCTU - HYPOTHETICAL 170	2316	Novel Protein sim. GBank			264906, 264762, 264687, 264769, 264689,
Novel Protein sim. GBank gil136406 dbj BAA11490 - (D79995) similar to pig tubulin-tyrosine ligase. [Homo saplens] Novel Protein sim. GBank gil2443342 dbj BAA22380 - UNCLASSIFIED (D88764) alpha 2 type I collagen [Rana catesbeiana] UNCLASSIFIED Novel Protein sim. GBank gil2443342 dbj BAA22380 - UNCLASSIFIED gil5503375 refilNP_001376.1 pDPYS - dihydropyrimidinase UNCLASSIFIED Novel Protein sim. GBank gil4589476 dbj BAA76766.1 - UNCLASSIFIED Novel Protein sim. GBank gil4589476 dbj BAA76766.1 - UNCLASSIFIED (AB023139) KIAA0922 protein [Homo sapiens] UNCLASSIFIED		gi 2496481 sp Q50724 Y09S_MYCTU - HYPOTHETICAL 87.3 KD PROTEIN CY78.27C			18108374, 35695855
1136406 dbj BAA11490 -	318				264687
In-tyrosine ligase. [Homo	232	Novel Protein sim. GBank gil			29331822, 29331824, 66714117, 29331826,
UNCLASSIFIED UNCLASSIFIED Iransport UNCLASSIFIED			-		60433356, 265018, 265019, 83373044, 18108385, 22279000, 22279002, 264563
UNCLASSIFIED UNCLASSIFIED transport UNCLASSIFIED	322				264369
UNCLASSIFIED transport UNCLASSIFIED	232		ח	NCLASSIFIED	264693
transport UNCLASSIFIED	232	=	ח	NCLASSIFIED	29331827, 264906
UNCLASSIFIED	2328	1	<u> </u>	ansport	18108398, 29331827, 29331828, 29146498,
UNCLASSIFIED		∓ਾ≡			29146499, 18108354, 21906768, 29148627, 21906769, 264693, 18108382, 18108385
	233(ח	INCLASSIFIED	264602, 264605, 264769, 18108370, 18108374, 264565
	233	Novel Protein sim. GBank gil4589476 dbj BAA76766.1 -			264488, 35696286, 22278999, 264259, 66714117 60432289, 35698052, 264905,
21906754, 33109954, 265017, 265018 264448, 264268, 264766, 264685, 350 13696855, 264558, 18108385, 60432		(Aboza 139) NAACOSZ protein (abrica sapiena)			56182435, 265006, 60433438, 264759.
264448, 264268, 264760, 264603, 33 34694848, 264288, 264760, 264888, 80432					21906754, 33109954, 265017, 265019,
					264448, 264268, 264768, 264663, 33636423, 13664885, 264558, 18108385, 60432113

1167	7006 2007 (220 222)	Marie Designation Court attended to the state of the stat			
	19905002 (2993, 2994)	(2555, 2554) Nover Protein Sim. Chank gitzbu433[dbj BAA23138] - (D76414) ppGpp hydrolase (Staphylococcus aureus)		kinase	264488
	88094678 (2335, 2336)			UNCLASSIFIED	264259, 29331827, 56182435, 60433438,
	11805403 (2337, 2338)			l	264684
_	21632244 (2339, 2340)				264602
1171	20434582 (2341, 2342)	20434582 (2341, 2342) Novel Protein sim. GBank gil2772914 (AF029249) -		UNCLASSIFIED	284556
1172	70610113 (2343 2344)	70610113 (2312 2311) National Bestin Co.			
	(2343, 2344)	Nover Protein sim. GBank gil4757846 ref NP_004317.1 pBCL9 - B-cell CLL/lymphoma		UNCLASSIFIED	55810764, 35696052, 264634, 264486
_	100001010000000000000000000000000000000	6			
?	50235713 (2345, 2345)	00233713 (2345, 2346) Novel Protein sim. GBank gi]2564053 dbj BAA22946 -			264508, 264906, 264907, 264909, 264591,
1174	20293077 (2347, 2348)	20293077 (2347, 2348) Novel Protein sim GBank giodingyx mort			264632, 264638, 264639
	((AL021958) mmsA (Mycobacterium tuberculosis)		denydrogenase	264600
1175	20711847 (2349, 2350)	20711847 (2349, 2350) Novel Protein sim. GBank	Ŀ	carboxylase	264601
		gi[118333 sp[P23234 DCIP_ENTCL - INDOLE-3-	Thiamine pyrophosphate enzymes		
		FINOVATE DECARBOATCASE (INDOLEPTRUVATE DECARBOXYLASE)			
1176	30252645 (2351, 2352)	80252845 (2351, 2352) Novel Protein sim. GBank gij1144520 (U34956) -	Contains protein domain (PF00586) - synthase	synthase	264509 264905 264593 264602 264605
		phosphoribosylformylglycinamidine synthase	AIR synthase related protein		10000 TOTAL
_		[Mycobacterium tuberculosis]			
1177	30064647 (2353, 2354)	80064647 (2353, 2354) Novel Protein sim. GBank	Contains protein domain (PF00106) - reductase		264605
_		gil119791 splP28643 FABG_CUPLA - 3-0XOACYL-[ACYL-			
		CARRIER PROTEINJ REDUCTASE PRECURSOR (3-			
_		KETOACYL-ACYL CARRIER PROTEIN REDUCTASE)			
<u> </u>	4128641 (2355, 2356)		Contains protein domain (PF00122) -	ATPase_associated	Contains protein domain (PF00122) - ATPase_associated 65274572, 18108398, 22278998, 22278999,
			E1-E2 ATPase		29331826, 264508, 264908, 264828,
		cholestasis 1, (progressive, Byler disease and benign			33657402, 33109954, 264769, 21906765,
		recurrent)			21906766, 21906768, 55811957, 33657023,
					264629, 55811576, 35696423, 264636,
					264556, 56182323, 60432113, 22279000,
1179 8	10055575 (2357, 2358)	80055575 (2357, 2358) Novel Protein sim. GBank gil2960090 emb CAA17988.1 -	Contains protein domain (PF00496) -	transport	264603
		(AL022121) dppA [Mycobacterium tuberculosis]			
1180	1794446 (2359, 2360)	11794446 (2359, 2360) Novel Protein sim. GBank gil2558614lemblCAA047871.	Contains protein domain (DE01220) sunthass		26.162.0
		(AJ001493) dehydroguinate dehydratase (Strentomyces	Debydronings a class II		000000
j	_	coelicolor)			
\neg	17946362 (2361, 2362)			UNC! ASSIFIED	65017
1182 8	1494264 (2363, 2364)	81494264 (2363, 2364) Novel Protein sim. GBank gi[5420387]emb CAB46679.1 -			265007, 265009, 264564, 264909, 264693
	10000 10001 10001	(AJZ43459) proteophosphoglycan [Leishmania major]			
201	78574044 (2365, 2366)				264689, 35696423, 264638, 18108385
_	233833 (2367, 2368)	ozobaso (zocz, zobó) Novel Protein sím. GBank gil4091877 (AF061331) - aipha Igalactosidase precursor (Saccharopolyspora erythraea)		UNCLASSIFIED	264602
1185 7	9491185 (2369, 2370)	79491185 (2369, 2370) Novel Protein sim. GBank gi[2129478 piri 551939 -		glycoprotein	263967
		Chimidse (EC 3.2.1.14) precursor - beet			

g	CT-00 1-001 C 1010COOL				
1				UNCLASSIFIED	264559
188		Novel Protein sim CBank ai		UNCLASSIFIED	29331825, 265017, 18108351
. 1		Invocational salts (AFU34325) - Invocational Process (AFU34325) - Invocational protein (Synechococcus PCC7002)		UNCLASSIFIED	264905, 264906
1189					264602
ol				I INC. ASSIEIED	26018
1191				UNCLASSIFIED	264600, 264605, 264764, 35695855, 264638,
1192	13521641 (2383, 2384)		Contains protein domain (PF01581) -		264886 264636
1193	11103584 (2385, 2386)		rwkramide related peptide family		
4	78893947 (2387, 2388)	1194 78893947 (2387, 2388) Novel Protein sim. GBank gil854065lemhlCAA583371.		UNCLASSIFIED	263978
1104	_	(X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	265007, 265008
s 1.	_	20443444 (2309, 2390) Novel Protein sim. GBank gi]1790277 (AE000459) - putative oxidoreductase (Escherichia coli)	0	UNCLASSIFIED	264605
1196	_				2646RQ
		11392317 (2393, 2394) Novel Protein sim. GBank	Contains protein domain (PF00571) - dehydrogenase	dehydrogenase	264594
		SILASTANDISPLANT SIMUNT, MY CLU - INCSINE-5- MONOPHOSPHATE DEHYDROGENASE (IMP DEHYDROGENASE) (IMPDH) (IMPD)	CBS domain		
ا ـــا	95290101 (2395, 2396)	1198 95290101 (2395, 2396)			264603
_	[81882011 (2397, 2398)	Novel Protein sim GRank			
- 1.		gil709525 sp P54673 P3K1_DICDI - PHOSPHATIDYLINOSITOL 3-KINASE 1 (PI3-KINASE) (PTDINS-3-KINASE) (PI3K)			264259, 264757, 33109954, 21906768
307	9848880 (2399, 2400)			UNCLASSIFIED	264910
5	80503751 (2401, 2402)	80503751 (2401, 2402) Novel Protein sim. GBank gilz4998771splP70645[BLMH_RAT - BLEOMYCIN HYDROLASE (BLM HYDROLASE) (BMH)		cathepsin	264766, 284769
	80082633 (2403, 2404)	1202 60082633 (2403, 2404) Novel Protein sim. GBank gi 606342 (U18997) - ORF_0622; reading frame open far upstream of start; possible frameshift, linking to previous ORF [Escherichia coli]		ribosomalprot	264600, 264558
1203	82125373 (2405, 2406)			UNCLASSIFIED	264768, 264769, 35695917, 264910, 264760, 264908, 264907, 264629, 264908, 264909
- 1	2010 1010100000000000000000000000000000				264766
		60303916 (2407, 2408) Novel Protein sim. GBank gi 2500728 sp Q59912 SECY_STRGB - PREPROTEIN TRANSLOCASE SECY SUBUNIT			264905, 264769, 264636
				UNCLASSIFIED	264566
1207	70841402 (2411, 2412)			UNCLASSIFIED	264556, 264557, 264558
1208	87755217 (2415, 2415)	Control October 100 Control October 10 Control Octo			29331824, 264909, 265021, 18108370
	(2410, 2410)	r 552 in (24 i5, 24 i5) Indover Protein Sim. GBank gilz645560 (AF027954) - BG-2. related ovarian killer protein [Rattus norvegicus]	Contains protein domain (PF00452) - apoptosis Apoptosis regulator proteins, Bcl-2 family	apoptosis	29331824, 29331825, 29331827, 265007, 264764, 264683, 264769, 264688, 264689

4200	70405747 12447 24401	4000 70405742 44401 1 - 1 D - 1 - 1 D - 1 - 1 D - 1 - 1 D - 1 - 1	, Caccordo		000700 100700
		10000175033381 SIII. ODBINA 11770338191P44398 XYLA_HAEIN - XYLOSE	Xylose isomerase	200	990407 , 100407
,	_	COMERAGE			
3	_			UNCLASSIFIED	264907, 264693
1211	94665655 (2421, 2422)	94665655 (2421, 2422) Novel Protein sim. GBank gil421095 pir S30688 - hypothetical protein 0246 - Escherichia coli		transferase	264591, 264592, 264595
1212	_	79167929 (2423, 2424) Novel Protein sim. GBank gij3880625[emb[CAB07858] -	Contains protein domain (PF01412) -		264689, 263967
		(293785) predicted using Genefinder; similar to RNA	Putative GTP-ase activating protein		
		recognition motif. (aka RRM, RBD, or RNP domain); cDNA	for Art		
		EST EMBL: T01682 comes from this gene; cDNA EST			
		EMBL:M75823 comes from this gene; cDNA EST			
		EMBL: D27559 comes from this ge			
1213		Novel Protein sim. GBank gi[226292]prf[[1505375A - vir		kinase	264909
		gene (Bordetella pertussis)			
1214	_	10144306 (2427, 2428) Novel Protein sim. GBank		UNCLASSIFIED	264908
		gi 5726285 gb AAD48396.1 AF12616 - (AF126162) HERV-H			
		LTR associating protein 2 [Homo sapiens]			
1215		80050106 (2429, 2430) Novel Protein sim. GBank gi[2326739]emb CAB10953 -		UNCLASSIFIED	265009, 264601, 264602, 264603, 33657109
	_	(Z98268) recN [Mycobacterium tuberculosis]			
1216	20438324 (2431, 2432)	20438324 (2431, 2432) Novel Protein sim. GBank		transferase	264604
		gij417329ispiP33038iMURA ENTCL - UDP-N-			
		ACETYLGLUCOSAMINE 1-			
		CARBOXYVINYLTRANSFERASE (ENOYLPYRUVATE			
		TRANSFERASE (110P.N.ACETY) GUICOSAMINE			
		ENO! PYRIVY! TRANSFERASE! (FPT)			
1217	95011344 (2433, 2434)			UNCLASSIFIED	264905 264907 264908 264591 264766
					264691 264693 264629 264630 264636
					264564
1218	110036RD (2435, 2436)	110936R0 (2435, 2436) Novel Protein sim Chank nitt 8054601dhill 4 Angologi.		dobydrocopaco	254501
2	100000 (2100, 2100)			ace in Ac	00000
		dehydrogenase GabD of E. coli (Bacillus subtilis)			
1219	91216252 (2437, 2438)				56181686, 29331822, 60432289, 264601, 264692, 264629
5	_	The Court of the C			Crosse double dorse source transcer
0221		81241324 (2438, 244U) Novel Protein sim. GBank gij424U313 db BAA74835.1 - /ABD37730 VIAA0043 Ecotois (Home conjoca)		oncogene	326443U7, 2645U3, 2649U9, 263U08, 263U19,
_		(suantendral protein jugana sabiens)			ZODUZU, DZB441DU, SSOD/UZS, ZB4693,
					33657182, 35695763, 264634, 22279000, 22279002, 264482
1221	83045055 (2441, 2442)	83045055 (2441, 2442) Novel Protein sim. GBank gil2143886lpirll152523 -		UNCLASSIFIED	264768, 265020, 264906
1222	20711865 (2443, 2444)	20711865 (2443, 2444) Novel Protein sim. GBank	Contains protein domain (PF00486) - phosphatase	phosphatase	264601
		gij730805jspijP39663jSPHR_SYNP7 - ALKALINE	Transcriptional regulatory protein, C	•	
_		PHOSPHATASE SYNTHESIS TRANSCRIPTIONAL	terminal		
		REGULATORY PROTEIN SPHR			
1223	11615647 (2445, 2446)				264593
1224	80432645 (2447, 2448)	80432645 (2447, 2448) Novel Protein sim. GBank	Contains protein domain (PF01472) - kinase	kinase	264593, 264600, 264601, 264603, 264605.
		gil1172627[spiP46546]PROB_CORGL - GLUTAMATE 5-	PUA domain		264768, 18108376, 264635, 18108387
		KINASE (GAMMA-GLUTAMYL KINASE) (GK)			

1225					
		800337418 (2454 2453) Novel Bodein ein Obert Britain			264768
	(201, 2405)	Inover Frutein sim. Gbank gijz 103030jembjCAB08836j - (295436) hypothetical protein Rv3644c (Mycobacterium juberculosis)		роlутегаѕе	264905, 264512, 264589
1227	79422138 (2453, 2454)	79422138 (2453, 2454) Novel Protein sim. GBank gi 1706768 sp P98133 FBN1_BOVIN - FIBRILLIN 1 PRECURSOR (MP340)		UNCLASSIFIED	264908, 264637, 264639
1228	79209027 (2455, 2456)	Novel Protein sim. GBank gi 1653901 db BAA18811 - (D90917) acriflavine resistance protein Synechocystls sp.)	Contains protein domain (PF00873) - Acr8/AcrD/AcrF family		264605, 264634
1229	94329135 (2457, 2458)			UNCLASSIFIED	87168474, 265011, 87168559, 264681, 264689, 264693, 65274620, 18108374
1230	80049357 (2459, 2460)	80049357 (2459, 2460) Novel Protein sim. GBank gil116230 sp P28598 CH60_BACSU - 60 KD CHAPERONIN (PROTEIN CPN60) (GROFI PROTEIN)	BACSU - 60 KD CHAPERONIN TCP-1/cpn60 chaperonin family PROTEIN	eph	264909, 264605, 18108388
1231	79843141 (2461, 2462)			INCI ASSIEIED	264008
1232	79853104 (2463, 2464)	79853104 (2463, 2464) Novel Protein sim. GBank gi[1215733 (U48718) - OphC [Agrobacterium tumefaciens]		transport	264909
1233	80255179 (2465, 2466)			UNCLASSIFIED	265017, 264564
		gij116298 sp P20730 CHHC_BOMMO - CHORION CLASS HIGH-CYSTEINE HCB PROTEIN 13 PRECURSOR (HC- B.13)			
1234	79242158 (2467, 2468)	79242158 (2467, 2468) Novel Protein sim. GBank ai7296711splP40280H2A MAI7F - HISTONE H2A	Contains protein domain (PF00125) - histone	histone	265008, 265010, 18108381
1235	79914423 (2469, 2470)		ALICUIGATION DIOSE DIOSE		264674 264762
	81927147 (2471, 2472)			UNCLASSIFIED	265018 55811150 264565 264757
1237	83371782 (2473, 2474)	83371782 (2473, 2474) Novel Protein sim. GBank gij3875133jemb CAA94750j -			264758, 264601, 264766, 264687, 18108372,
		(L7.07.50) similar to actin binding domain; CDNA EST EMBL:100093 comes from this gene; CDNA EST EMBL:03443 comes from this gene; CDNA EST EMBL:037508 comes from this gene; CDNA EST EMBL:084247 comes from this gene; CDNA EST EMBL:			264555, 264559
1238	87411577 (2475, 2476)	1===	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	glycoprotein	264259, 29331822, 29331824, 35696052, 264508, 264906, 52644045, 52646317, 264288, 264769, 264693, 264632, 264634,
1239	82197449 (2477, 2478)	82197449 (2477, 2478) Novel Protein sim. GBank gil4007990 gbJpAC95339 - AF084363) DOK protein Mus musculus		oncogene	264558, 87168518, 284563 264509, 264511, 264759, 264760, 264764,
1240	80497259 (2479, 2480)	80497259 (2479, 2480) Novel Protein sim. GBank gij1176192 sp P45420 YHCD_ECOLI - HYPOTHETICAL OUTER MEMBRANE USHER PROTEIN IN GLTF-NANT INTERGENIC REGION PRECURSOR			264769
1241		1241 80020711 (2481, 2482) Novel Protein sim. GBank gi 121383 sp P19904 GLNA_VIBAL - GLUTAMINE SYNTHETASE (GLUTAMATE-AMMONIA LIGASE)	Contains protein domain (PF00120) - UNCLASSIFIED Glutamine synthetase	UNCLASSIFIED	264601, 264604, 264638
1242	1242 79775890 (2483, 2484)				264906, 264907, 264908, 264634

5	(3010, 3010)	AND TOTTOLED DADE DADES IN TOTOL CONTRACTOR CONTRACTOR			
?	(2012, 2450 (2463, 2460)	(ALO31124) branched-chain amino acid aminotransferase		UNCLASSIFIED	18108374, 35695917, 35695855, 265009, 264508, 264909
		[Streptomyces coelicolor]			
1244	10284821 (2487, 2488)	10284821 (2487, 2488) Novel Protein sim. GBank gil2970646 (AF051945) - Xin		UNCLASSIFIED	264691
		[Mus musculus]			
1245	80437103 (2489, 2490)	80437103 (2489, 2490) Novel Protein sim. GBank gi[4586338 dbj]BAA76357.1 -	-	oxidase	264768
		[Pseudomonas putida]	cylocarome c and cultor exidase polypeptide I		
1248	80059321 (2491, 2492)			UNCLASSIFIED	264604, 264636, 264557, 264564
		(AL031541) putative phenylalanyl-tRNA synthetase beta		-	
		chain [Streptomyces coelicolor]			
1247	80064831 (2493, 2494)	Novel Protein sim. GBank gi 2621684 (AE000842) -			264758, 264605, 264639
	_	adhesion protein [Methanobacterium thermoautotrophicum]			
1248	_	88070353 (2495, 2496) Novel Protein sim. GBank	Contains protein domain (PF00316) - UNCLASSIFIED	UNCLASSIFIED	18108392, 264259, 29331826, 264106,
		gij1352403 spjP09467 F16P_HUMAN - FRUCTOSE-1,6-	Fructose-1-6-bisphosphatase		264508, 264907, 264828, 265009, 60433356,
		BISPHOSPHATASE (D-FRUCTOSE-1,6-BISPHOSPHATE			264757, 264758, 21906754, 265010, 265011.
		1-PHOSPHOHYDROLASE) (FBPASE)			265018, 265019, 264760, 18108351.
					18108354, 265021, 18108376, 18108377, 264630, 18108385
1249	80056657 (2497, 2498)	80056657 (2497, 2498) Novel Protein sim. GBank gil2791407 lembiCAA160011 -		transport	264908 265010 264600 264603 264691
		(At 021184) hypothetical protein By1473 (Mycohacterium			18108376
		[treezerosis]			
1250	12694385 (2499 2500)	12694385 (2499, 2500) Novel Protein sim GBank		INCI ASSIFIED	264689
		METHYLADENINE GLYCOSIDASE 1/3-METHYLADENINE.			
		ONA CI VODEVI ASE I CONSTITUTO LA LACITA			
1251	79850448 (2501, 2502)	79850448 (2501, 2502)			264909
	79458087 (2503 2504)			I INC. A COLETED	264683 263076
1367	00050404 (2505, 2505)	Married Description Consult		חווייים אים	201000, 201000 10100010
	80050121 (2505, 2506)			glycoprotein	264600, 264603, 181083/6
		gi5670176[gb]AAD46616.1[AF16131 - (AF161317) NRAMP			
		manganese transport protein MntA (Salmonella tvohimirrium)			
1254	87716767 (2507, 2508)	87716767 (2507, 2508) Novel Protein sim. GBank gil103160lpirllS22126 - finger		UNCLASSIFIED	35696286 264910 264764 264688
		protein unkempt - fruit fly (Drosophila metanogaster)			21906767, 55811957, 264692, 264556,
					264639
1255					264636
1256		87889508 (2511, 2512) Novel Protein sim. GBank gi[2995353 emb CAA04608.1 -		UNCLASSIFIED	60432289, 264600, 264605, 264764, 264687,
		(AJ001206) pep2 [Streptomyces coelicolar]			264769, 264689, 27486265, 18108374,
					18108376
1257	80201435 (2513, 2514)	80201435 (2513, 2514) Novel Protein sim. GBank gi 3193306 (AF069300) -		UNCLASSIFIED	264094, 265019
		contains similarity to Arabidopsis membrane-associated salt			
		inducible-like protein (GB:AL021637) [Arabidopsis thaliana]			
1258	20708150 (2515, 2516)			INCLASSIFIED	264602 263978
1250	B0186012 (2517 2518)			INCI ASSIEIED	DEAGUE DEAGAR DEAGUR
1000	90004606 (2640, 2620)				SCASSOL, SCATTS, SCASSOS
8	00004000 (2318, 2320)			UNCLASSIFIED	204034, 204038

1261	87412802 (2521, 252	87412802 (2521, 2522) Novel Protein sim. GBank gi[5689511 dbj BAA83039.1 -	Contains protein domain (PF01699) - Cadherin	cadherin	29331824, 264906, 264909, 264768, 264769,
			Sodium/calcium exchanger protein		264689, 264693, 264639, 18108384, 264563
1262	13504589 (2523, 252)	13504589 (2523, 2524) Novel Protein sim. GBank gi[95100 pir] S21334 - hypothetical protein 4 - Agrobacterium tumefaciens		UNCLASSIFIED	264634
1263		(9)			264602
1264		80083396 (2527, 2528) Novel Protein sim. GBank gij3550958 (AF004840) - CDO [Rattus norvegicus]		struct	264634
1265		(0)		UNCLASSIFIED	264563
1266	Ī			UNCLASSIFIED	264766, 264636, 264638, 264567
1267		Novel Protein sim. GBank gi 1085002 pir S55056 - mitochondrial carrier protein DIF-1 homolog - Caenorhabditis elegans	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins		264259, 21906754, 264369
1268		88178473 (2535, 2536) Novel Protein sim. GBank gil488645 emb CAB43370.1 - (AL050269) hypothetical protein [Homo saplens]	Contains protein domain (PF00583) - UNCLASSMED Acetyltransferase (GNAT) family	UNCLASSIMED	18108398, 22278995, 56994075, 60424269, 29331827, 264109, 264512, 265007, 265008, 265009, 264595, 33109954, 33657084, 87168559, 264600, 265018, 265019, 264369, 264688, 21906767, 265020, 52644150, 264691, 33657023, 33657349, 18108374, 264566, 18108385, 60432113, 22279002, 264486
1269		79821946 (2537, 2538) Novel Protein sim. GBank gij3334791 lemb CAA19939 - (AL031107) hypothetical protein SC5A7.10c [Streptomyces coelicolor]		UNCLASSIFIED	264508, 264905, 264906, 264687, 264693
1270		80031420 (2539, 2540) Novel Protein sim. GBank gi 2851634 sp Q50591 Y0D1_MYCTU - HYPOTHETICAL 50.0 KD PROTEIN CY1A11.01	Contains protein domain (PF01574) - dehydrogenase IMP dehydrogenase / GMP reductase N terminus	dehydrogenase	265010, 264601
1271	79840499 (2541, 2542)			ATPase_associated 35696052, 264908	35696052, 264908
1272	_	(4)			264686, 264689
1273		80220315 (2545, 2546) Novel Protein sim. GBank gi 1655665 emb CAB03731 - (Z81368) hypothetical protein Rv2395 [Mycobacterium tuberculosis]		UNCLASSIFIED	264509, 264639
1274	ĺ	(81)		UNCLASSIFIED	264905, 264908, 264909, 264769
1275		20730763 (2549, 2550) Novel Protein sim. GBank gij123726jspjP10413jHTPG_ECOLI - HEAT SHOCK PROTEIN HTPG (HIGH TEMPERATURE PROTEIN G) (HEAT SHOCK PROTEIN C62.5)	Contains protein domain (PF00183) - eph Hsp90 protein	ph	264602
1276	_	21148644 (2551, 2552) Novel Protein sim. GBank gil2129478 pir S51939 - chitinase (EC 3.2.1.14) precursor - beet		UNCLASSIFIED	264369
1277	_	(3)		UNCLASSIFIED	264556
1278		11088365 (2555, 2556) Novel Protein sim. GBank gi 1175473 sp P44555 YAAJ_HAEIN - HYPOTHETICAL PROTEIN HI0183		l	264603
1279		21658756 (2557, 2558) Novel Protein sim. GBank gi 1929513 (U64318) - ATP synthase subunit beta [Moorella thermoacetica]		synthase	264605
1280	_	79310959 (2559, 2560) Novel Protein sim. GBank gil4938504 emb CAB43862.1 - (AL078465) putative protein [Arabidopsis thaliana]		struct	263976

Parizage (2561, 2562) Nover Protein sim. GBank gil 136501 (139548) - surface Contains protein domain (PF00047) - protein MCA-22 Battless novegeues Immunoglobulin domain (PF00047) - protein MCA-22 Battless novegeues Immunoglobulin domain (PF00047) - protein MCA-22 Battless novegeues Immunoglobulin domain (PF00163) - 2046505 (2563, 2564) Nover Protein sim. GBank gil 328190 (AF0266) - L59B Contains protein domain (PF00183) - [A87330 (2563, 2565) Nover Protein sim. GBank gil 32810 (AF0266) - L59B Contains protein domain (PF00183) - [A87330 (2564, 2571, 2572) Nover Protein sim. GBank gil 32820 (AF0266) - L59B Contains protein domain (PF00183) - [A87330 (2565, 2565) Nover Protein sim. GBank gil 32820 (AF088815) Entities protein domain (PF00183) - [A87330 (2575, 2572) Nover Protein sim. GBank gil 32820 (AF088815) Entities protein domain (PF00183) - [A873316 (2575, 2575) Nover Protein sim. GBank gil 32820 (AF088815) Entite	SSIFIED 29331825, 29331828, 264766, 83373044	UNCLASSIFIED 265008	UNCLASSIFIED 264605	264604	264766, 264689, 263967	se 264605, 264639	UNCLASSIFIED 265011, 264602, 264766, 284687, 264769, 264689, 18108370, 264636, 18108385, 264563		UNCLASSIFIED 264637			ptfactor 264092, 264259, 29331822, 29331824, 264508, 264909, 264909, 264512, 265008, 265009, 264090, 264369, 264288, 284686, 264768, 264693, 18108374, 264632, 264639, 83373044, 22279002, 264482, 264663	SFED		genase 264689
87337898 (2561, 2562) [Novel Protein sim. GBank gil1136501 (U39546) - surface protein MCA-32 [Rattus novegicus] oncogene AF4 [Mus musculus] (20468305 (2565, 2566) [Novel Protein sim. GBank gil328190 (AF074266) - proto-consequence AF4 [Mus musculus] (AF07632 [Mycobacterium tuberculosis] (292770) hypothetical protein Rw01532 [Mycobacterium tuberculosis] (292770) hypothetical protein Rw01532 [Mycobacterium tuberculosis] (AF086325 (2567, 2566) Novel Protein sim. GBank gil3929022 (AF057696) - L59B [Haemophilus ducreyi] (AF08812 (2571, 2572) [Movel Protein sim. GBank gil2078004 emb[CA808451] - (295207) gorld [Mycobacterium tuberculosis] (2573, 2574) [Movel Protein sim. GBank gil2078004 emb[CA808451] - (295207) gorld [Mycobacterium tuberculosis] (2575, 2578) Novel Protein sim. GBank gil2078074 emb[CA808305] - (29770) Movel Protein sim. GBank gil2072674 emb[CA808305] - (29770) Movel Protein sim. GBank gil2072674 emb[CA808305] - (29712316 (2581, 2582) Novel Protein sim. GBank gil2072674 emb[CA808305] - (295120) mil [Mycobacterium tuberculosis] (2583, 2584) Novel Protein sim. GBank gil2072674 emb[CA808305] - (295120) mil [Mus musculus] (2585, 2588) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (2588, 2588) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (2588, 2588) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (1688685) (2588, 2589) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (1688685) (2588, 2589) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (1688685) (2588, 2589) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.3) alpha subunit (1688685) (2588, 2589) Novel Protein sim. GBank gil20729173 pin [64453 - cardoacetale decarboxylase (EC 4.1.1.174) alpha subunit (1688685) (2588) (1688685) (2588) (168868	Contains protein domain (PF00047) - UNCLA		UNCLA		Contains protein domain (PF00183) - eph Hsp90 protein	reductase	UNCLA	Contains protein domain (PF00386) - collager C1q domain	UNCLAS	Contains protein domain (PF00001) - tm7 7 transmembrane receptor rhodopsin family)	Contains protein domain (PF00271) - helicase Helicases conserved C-terminal Jomain	Contains protein domain (PF01530) - transcrip Zinc finger, C2HC type	UNCLAS	biotinde	dehydrogenase
	Novel Protein sim. GBank gij1136501 (U39546) - surface protein MCA-32 [Rattus novveoicus]	Novel Protein sim. GBank gij3328190 (AF074266) - proto- oncogene AF4 [Mus musculus]	Novel Protein sim. GBank gij3261721 emb CAB07057 - (292770) hypothetical protein Rv0153c [Mycobacterium tuberculosis]	Novel Protein sim. GBank gij3929022 (AF057696) - LspB [Haemophilus ducreyi]	Novel Protein sim. GBank gil417154 sp P33126 HS82_ORYSA - HEAT SHOCK PROTEIN 82	Novel Protein sim. GBank gi 2078004 emb CAB08451 - (295207) gorA [Mycobacterium tuberculosis]		Novel Protein sim. GBank gi[5353510]gb AAD42161.1 AF08891 - (AF088916) emilin precursor [Homo sapiens]		Novel Protein sim. GBank gil 169995jspjP46023jGPCR_LYMST - G-PROTEIN COUPLED RECEPTOR GRL101 PRECURSOR	gij2072674jemb CAB08305 - rium (uberculosis)	gi 1835755 (U86338) - zinc finger lius]		Novel Protein sim. GBank gi[2129173]pir F64453 - oxaloacetate decarboxytase (EC 4.1.1.3) atpha subunit - Methanococcus Jannaschii	Novel Protein sim. GBank gi[5441779]emb CAB46803.1 - (AL096811) putative alcohol dehydrogenase (zinc-binding)
1281 1283 1283 1285 1286 1286 1289 1290 1293 1293 1294 1294	1281 94323988 (2561, 2562)									19526027 (2579, 2580)	80470268 (2581, 2582)	94723316 (2583, 2584)	_	82125908 (2587, 2588)	11686851 (2589, 2590)

430	4206 144002004 (2504 2502)				
202	_			UNCLASSIFIED	264591, 264639
2	_		Contains protein domain (PF01367) - polymerase	polymerase	264693
		gll+962191 gb AAU356865.1 AE00180 - (AE001805) DNA- directed DNA polymerase I [Thermotoga maritima]	5-3' exonuclease		
1298		94239506 (2595, 2596) Novel Protein sim. GBank gi 1943770 (U97191) - F53F10.1 [gene product [Caenorhabditis elegans]		struct	18108348, 265017
138	\neg	80255378 (2597, 2598)			264488, 264906, 264909, 22279002, 264566
300		Novel Protein sim. GBank gij3445181 (AC005498) - R31655_2 [Homo sapiens]	Contains protein domain (PF01352) - transcriptfactor KRAB box	transcriptfactor	264605
<u> </u>		17939614 (2601, 2602) Novel Protein sim. GBank gil4062973 dbj BAA36204.1 - (AB017138) alpha subunit of malonate decarboxylase Pseudomonas putida}		UNCLASSIFIED	264906
1302		_			85658542 265020
1303	3 9684121 (2605, 2606)				264908
ğ	_			UNCLASSIFIED	264508
8					264566
2 00		13069230 (2611, 2612) Novel Protein sim. GBank gij3242273 emb CAB07017 - (292669) hypothetical protein Rv0236c Mycobacterium tuberculosis}		UNCLASSIFIED	264636
1307				UNCLASSIFIED	264907, 264592, 264764
13 <u>0</u>		21426814 (2615, 2616) Novel Protein sim. GBank gi 1502421 (U59433) - 3-ketoacyl acyl carrier protein reductase [Bacillus subitiis]	- 3-ketoacyl Contains protein domain (PF00516) - reductase Envelope glycoprotein GP120	reductase	264555
1309		79263011 (2617, 2618) Novel Protein sim. GBank gil95819 pir S16298 - ferric enterobactin transport protein fepC - Escherichia coli		transport	264906, 18108354
1310				UNCLASSIFIED	264605
1311	1 87613142 (2621, 2622)				35696286, 29331827, 264908, 265008, 264764, 264766, 264686, 21906767, 21906769, 3565691, 264693
1312		88061720 (2623, 2624) Novel Protein sim. GBank gil4455118 gb AAD21084 - (AF125158) zinc finger DNA binding protein 99 [Homo sapiens]	Contains protein domain (PF00096) - dna_rna_bind Zinc finger, C2H2 type	dna_rna_bind	22278995, 22278998, 22278999, 264905, 264905, 264906, 265011, 265017, 265019, 264687, 265026, 265021, 33657023,
1313	_	91225458 (2625, 2626) Novel Protein sim. GBank	Contains protein domain (PE00886) - ribosomalprot	ribosomalorot	2227 3002, 204304 2227 8996 2227 8999 264259 2028 1099
		gl/4929733 gb AAD34127.1 AF15189 - (AF151890) CGI-132 Ribosomal protein S16	Ribosomal protein S16		29146498, 264508, 264908, 66712502,
		protein [Homo sapiens]			60433356, 60433438, 265011, 265017, 264683, 264288, 21906765, 21906767
					29148627, 21906768, 35695917, 265021,
					33657023, 33657109, 18108370, 18108377, 35695855, 60432113, 22279000, 264563.
1314					264693
1315		84357192 (2629, 2630) Novel Protein sim. GBank gil2589223 (AF026565) - ring finger protein [Mus musculus]	Contains protein domain (PF00097) - interleukinrecept Zinc finger, C3HC4 type (RING finner)	interleukinrecept	264691
			inger)		

UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED START domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type Start domain (PF01852) - synthase Contains protein domain (PF01852) - synthase Thrombospondin type 1 domain Thrombospondin type 1 domain START domain UNCLASSIFIED UNCLASSIFIED	2632	1316 [95361609 (2631, 2632) Novel Protein sim. GBank gil5689407 dbilBAA82987.11-	4	kinase	56182575, 56181686, 20281171, 29331822.
1/4F13491 - (AF134918)	₹ V	028958) KIAA1035 protein [Homo sapiens]			29331824, 60424269, 29331825, 35696052, 52644045, 264591, 60432229, 265018
14F13491 - (AF134918)					265019, 55811150, 56181562, 21906765,
14F13491 - (AF134918)					21906767, 21906768, 35695917, 60170615,
14F13491 - (AF134918)					33657023, 65274620, 33657109, 35695763,
1/6F13491 - (AF134918)					35695855, 18108387, 87168518, 60432113, 22279002, 264564
14F13491 - (4F134918)	ž	yvel Protein sim. GBank		UNCLASSIFIED	264093, 264906, 264909, 264369, 264684
UNCLASSIFIED UNCL	<u> </u>	4836757[gbtAAD30541.1jAF13491 - (AF134918) maphorin subclass 4 member G [Mus musculus]			
lum bicolor] lum bicolor] lum bicolor] lum bicolor] leain [Homo saplens] gij4240183[dbi]BAA74870.1] - Contains protein domain (PF00651) - transcriptfactor rotein [Homo saplens] gij4886505[emb]CAB4377.1] - Contains protein domain (PF00096) - dna_ma_bind rotein [Homo saplens] gij5262591[emb]CAB45736.1] - Zinc finger, C2H2 type gij5265591[elbi]BAA82407.1] - Sinc finger, C2H2 type thanolamine N-methyltransferase gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease hibitors; most similar to tissue Thrombospondin type 1 domain gij1397275 (U61947) - C06G3.8 gij1397275 (U61947) - C06G3.8 gij1397275 (U61947) - C06G3.8 gij1397275 (U61947) - C06G3.8	Ĭ	ovel Protein sim. GBank		UNCLASSIFIED	18108392, 18108348, 265011, 265017,
gij4240183jdbjjBAA74870.1 - tein [Homo sapiens] gij4886505jembjCAB43377.1 - gij4886505jembjCAB45738.1 - gij5262591jembjCAB45738.1 - gij5262591jembjCAB45738.1 - gij5355595jembjCAB45738.1 - gij5359516jdbjjBAA82407.1 - thanolamine N-methyltransferase gij5459516jdbjjBAA82407.1 - thanolamine N-methyltransferase gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease gij3294501 (U64857) - similar to Gontains protein domain gij3294501 (U64857) - similar to Gontains protein domain gij3394501 (U64857) - Similar to Gontains protein domain gij3394501 (U64857) - Similar to Gontains protein domain gij1397275 (U61947) - C06G3.8 gij1397275 (U61947) - C06G3.8	5 6	4680204 gb AAD27567.1 AF11417 - (AF114171)			18108359, 18108362, 56182323, 18108385,
tein [Homo sapiens] gil4886505[emb]CAB43377.11 - Contains protein domain (PF00651) - Iranscriptfactor rotein [Homo sapiens] gil5262591[emb]CAB45736.11 - Contains protein domain (PF00096) - dna_ma_bind gil5262591[emb]CAB45736.11 - Zinc finger, C2H2 type gil5459516[db][BAA62407.1] - IpCPBP - goodpasture antigen- START domain Contains protein domain (PF01852) - START domain Gil3294501 (U64857) - similar to Thrombospondin type 1 domain FUNCLASSIFIED gil3394501 (U64857) - co6G3.8 gil337275 (U61947) - Co6G3.8 gil337275 (U61947) - C06G3.8	_				264908, 264909, 265006, 265008, 264592.
gil4240183[dbj BAA74870.1] - tein [Homo sapiens] gil4886505[emb]CAB43377.1] - totein [Homo sapiens] gil5262591[emb]CAB45736.1] - totein [Homo sapiens] gil5262591[emb]CAB4573.1] - totein [Homo sapiens] gil5262591[emb]CAB4573.1] - totein [Homo sapiens] gil5262591[emb]CAB4573.1] - totein [Homo sapiens] gil5262591[emb]CAB482407.1] - totein [Homo sapiens] gil5262591[emb]CAB4825] - totein [Homo sapiens] gil5262591[emb]CAB4825] - totein [Homo sapiens] gil5262591[emb]CAB4827] - totein [Hom					265019, 264766, 56181562, 18108368,
gil4240183 dbj BA474870.1 - tein [Homo sapiens] gil4886505 emblCAB43377.1 - Totein [Homo sapiens] gil5262591 emblCAB45736.1 - Totein [Homo sapiens] gil5262591 emblCAB4573.1 - Tinc finger, C2H2 type synthase synthase gil5294501 (U64857) - similar to Thrombospondin type 1 domain protease Thrombospondin type 1 domain gil1397275 (U61947) - C06G3.8 gil1397275 (U61947) - C06G3.8 gil1397275 (U61947) - C06G3.8	_				264628, 264629, 18108377, 264636
Nein [Homo sapiens] Contains protein domain (PF00651) - transcriptfactor gil4886505[emb]CAB43377.1 - BTB/POZ domain Intein [Homo sapiens] gil5262591[emb]CAB45736.1 - Contains protein domain (PF00096) - dna_ma_bind gil5262591[emb]CAB482407.1 - Zinc finger, C2H2 type gil5459516[db][BA482407.1 - Zinc finger, C2H2 type gil5459516[db][BA482407.1 - Zinc finger, C2H2 type gil529516[db][BA482407.1 - Zinc finger, C2H2 type gil3294501 (U64857) - similar to Contains protein domain (PF00090) - protease hhibitors; most similar to tissue ecursor [Caenorhabditis elegans] gil1397275 (U61947) - C06G3.8 gil1397275 (U61947) - C06G3.8	Ž			UNCLASSIFIED	35696286, 55812038, 265018, 21906768,
gil4886505[emb[CAB43377.1] - Contains protein domain (PF00651) - transcriptfactor rotein [Homo sapiens] BTB/POZ domain Sinthase Synthase Synthase START domain Bij3294501 (U64857) - similar to Contains protein domain (PF00990) - protease Thrombospondin type 1 domain Bij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease Thrombospondin type 1 domain Bij3294501 (U64857) - co663.8 BIJ397275 (U61947) - C0663.8 BIII397275 (U61947) - C0663.8	ತಿ				265020, 263978, 22279002
rotein (Homo sapiens) gi[5262591jemb]CAB45736.1] - Contains protein domain (PF00096) - dna_ma_bind rotein [Homo sapiens] gi[5262591jemb]CAB45736.1] - Zinc finger, C2H2 type gi[5459516jdb][BAA82407.1] - Zinc finger, C2H2 type synthase gi[5459516jdb][BAA82407.1] - Contains protein domain (PF01852) - Similar to Contains protein domain (PF00090) - protease hhibitors; most similar to tissue ecursor [Caenorhabditis elegans] gi[1397275 (U61947) - C06G3.8 gi[1397275 (U61947) - C06G3.8 gi[1397275 (U61947) - C06G3.8	<u> </u>		Contains protein domain (PF00651) - 1	transcriptfactor	60432049, 29331828, 264907, 264908,
gi[5262591[emb]CAB45736.1] - Contains protein domain (PF00096) - dna_ma_bind rotein [Homo sapiens] Zinc finger, C2H2 type synthase gi[5459516]db]BAA82407.1] - Contains protein domain (PF01852) - Contains protein domain (PF01852) - START domain (PF00090) - protease protein domain (PF00090) -	≥		BIB/PO2 domain		204909, 204910, 33012030, 204001, 204702, 264764, 264766, 264768, 264769, 264628
gi[5262591[emb]CAB45736.1] - Contains protein domain (PF00096) - dna_ma_bind rotein [Homo sapiens] Zinc finger, C2H2 type gi[5459516]dbj[BAA82407.1] - Zinc finger, C2H2 type synthase thanolamine N-methyltransferase Contains protein domain (PF01852) - START domain gi[3294501 (U64857) - similar to Contains protein domain (PF00090) - protease hhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gi[1397275 (U61947) - C06G3.8 titis elegans]					18108374, 264634, 264635, 18108385
rotein [Homo sapiens] Zinc finger, C2H2 type synthase gil5459516[dbj BA482407.1] - thanolamine N-methyltransferase Contains protein domain (PF01852) - START domain Gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease hibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 titis elegans]	烏	gi 5262591 emb CAB45736.1 -	Contains protein domain (PF00096) - (dna_ma_bind	29331828, 264908, 265020, 33657023,
gil5459516[dbj BA482407.1] - thanolamine N-methyltransferase Contains protein domain (PF01852) - START domain Gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease Inhibitors; most similar to tissue Thrombospondin type 1 domain gij37275 (U61947) - C06G3.8 gil1397275 (U61947) - C06G3.8 titis elegans]		rotein (Homo sapiens)	Zinc finger, C2H2 type		264693, 264404
thanotamine N-methyttransferase Contains protein domain (PF01852) - Contains protein domain (PF01852) - START domain gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor (Caenorhabditis elegans) gij1397275 (U61947) - C06G3.8 UNCLASSIFIED	=	gi 5459516 db BAA82407.1 -		synthase	65274572, 56994075, 264259, 29331822,
Contains protein domain (PF01852) - START domain Gondains protein domain (PF01852) - START domain gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 titis elegans]		AB029821) phosphatidylethanolamine N-methyltransferase			29331827, 264104, 56182435, 87168474,
Contains protein domain (PF01852) - Contains protein domain (PF01852) - START domain gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 UNCLASSIFIED	=	Homo sapiens]			18108351, 264288, 21906766, 21906767,
Contains protein domain (PF01852) - START domain START domain SI3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gi[1397275 (U61947) - C06G3.8 UNCLASSIFIED titls elegans]					35695917, 265020, 264693, 65274791, seresson reposta
1.1 pGPBP - goodpasture antigen-START domain gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 titis elegans]	╁		Contains protein domain (PF01852) -		22278996, 22278998, 29331828, 264905,
gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 UNCLASSIFIED titls elegans]		.1 pGPBP - goodpasture antigen-	START domain		264907, 29331830, 264908, 264510, 265008,
gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 UNCLASSIFIED titls elegans]	_	pinding protein			264595, 264759, 21906754, 265018, 264288,
gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain ecursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 UNCLASSIFIED titls elegans]					264768, 264769, 21906768, 265022,
gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease nhibitors; most similar to tissue Thrombospondin type 1 domain secursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 UNCLASSIFIED titls elegans]	_				18108376, 264631, 264632, 264634, 264636,
gij3294501 (U64857) - similar to Contains protein domain (PF00090) - protease hhibitors; most similar to tissue Thrombospondin type 1 domain scursor [Caenorhabditis elegans] gij1397275 (U61947) - C06G3.8 titis elegans]	_				264638, 264563, 264564, 264565, 264566
hibitors; most similar to tissue Thrombospondin type 1 domain cursor [Caenorhabditis elegans] gil1397275 (U61947) - C06G3.8 UNCLASSIFIED tits elegans]	들	gi 3294501 (U64857) - similar to	Contains protein domain (PF00090) -	protease	35696286, 264905, 264906, 264907, 264908,
ecursor [Caenorhabditis elegans] gi[1397275 (U61947) - C06G3.8 litis elegans]	Ξ.		Thrombospondin type 1 domain		264909, 264910, 264593, 33657402, 264758,
gi 1397275 (U61947) - C06G3.8 Itis elegans]	4-	actor pathway inhibitor precursor (Caenorhabditis elegans)			85658542, 264760, 264768, 264769, 264691, 35696423
gene product (Caenorhabditis elegans)	ऻ ≂	Novel Protein sim. GBank gi[1397275 (U61947) - C06G3.8		UNCLASSIFIED	264259, 66712502, 264682, 264683, 264635
	- 1	gene product [Caenorhabditis elegans]			

1327	95322897 (2653, 2654)	95322897 (2653, 2654) Novel Prolein sim. GBank	Contains protein domain (PF00279) - UNCLASSIFIED		18108398, 22278996, 22278997, 22278999.
		BITZG63Z[SP]P39169[ALOZ_HUMAN - IIII ALU SUBFAMILY	HOWAN - IIII ALO SOBFAMILY Piant lipid transfer protein family		264091, 264259, 29331822, 29331824, 2633482£ 2633482£ 26334837 2633483
			,		264105, 264905, 56182435, 264112, 265008.
					265009, 21906754, 265010, 265011, 265017,
					265019, 264681, 264448, 264764, 264684,
					264288, 264685, 264768, 264688, 21906767,
					21906769, 29148629, 265020, 264690,
					264691, 264692, 264693, 263967, 33657109,
					33657182, 27486262, 33657349, 18108370,
					18108374, 55810764, 35695855, 264634,
					56182323, 83373044, 87168518, 60432113,
T	_			T	22279000, 22279002, 264563, 264567
1328	87753493 (2655, 2656)			UNCLASSIFIED	264488, 22278997, 29331826, 264595,
					18108351, 264766, 22279002, 264482, 264567
1329		87755276 (2657, 2658) Novel Protein sim GBank		LINCI ASSIFIED	22278996 29331827 264684 264692
		gil4678224[gblAAD26969,1]AC00713 - (AC007135)		_	33657109
		unknown protein (Arabidopsis thaliana)			
1330		Novel Protein sim. GBank gil437310 (L23504) - nodulin		UNCLASSIFIED	264259, 29331825, 264512, 265019, 265021,
		[Medicago truncatula]			264555, 264558, 56526486
1331		Novel Protein sim. GBank gil4589586 dbj BAA76815.1 -		UNCLASSIFIED	264259, 29331826, 29331827, 35696052,
		(AB023188) KIAA0971 protein [Homo sapiens]			29331828, 60170831, 264448, 264686,
					21906765, 55811957, 265020, 33657023,
					33657109, 263973, 55811576, 35696423,
T					35695855, 56182323
1332		94845937 (2663, 2664) Novel Protein sim. GBank gi 5459516 db BAA82407.1 -		synthase	65274572, 22278996, 56994075, 22278999,
		(AB029821) phosphatidylethanolamine N-methyltransferase			60432049, 264259, 29331822, 29331826,
		[Homo sapiens]			60432289, 29331827, 35696052, 52644045,
					56182435, 264510, 21906754, 87168559,
					265018, 265019, 264448, 264288, 264369,
					264686, 21906765, 21906766, 21906767,
					21906768, 265020, 265021, 52644150,
					33657023, 56182323, 18108387, 60432113,
					22279002
1333		88098476 (2665, 2666) Novel Protein sim. GBank gil5689527 dbj BAA83047.1 -		UNCLASSIFIED	60432289, 66712502, 264591, 60433356,
		(Abozeo 10) NiAA 1095 protein (Momo sapiens)			60433436, 53812036, 263010, 264639,
_	_				56526486
		87592388 (2667, 2668) Novel Protein sim. GBank gi 2662536 (AF036685) - Similar to protein-tyrosine phosphatase [Caenorhabditis elegans]		phosphatase	264905
1335	87644798 (2669, 2670)	87644798 (2669, 2670) Novel Protein sim. GBank gil4240285 db BAA74921.1 -	Contains protein domain (PF00643) - UNCLASSIFIED	Г	22278998, 22278999, 29331827, 264509.
	-	(AB020705) KIAA0898 protein [Homo sapiens]	B-box zinc finger.		264511, 265007, 265008, 265009, 60433438,
					21906754, 87168559, 265017, 264288,
					21906765, 21906767, 21906768, 21906769,
					265020, 33657109, 27486264, 18108374,
					264556, 264638, 264557, 60170394, 264559,
٦					18108385, 264563

264509, 264905, 264512, 264764, 264693, 264635, 264637	56182575, 56994075, 22278998, 22278999, 264092, 264259, 60432289, 29331826, 264006, 264908, 264102, 265008, 265009, 264112, 265008, 265009, 60433356, 55812038, 33657084, 265011, 265017, 265018, 265019, 264682, 264448, 264683, 264369, 264689, 21906766, 21906769, 265009, 264691, 27486261, 20281069, 18108379, 55811578, 35695855, 56182323, 60432113, 22279002, 264567	29331822, 265010, 264288, 264689, 18108370, 35695855	35696052, 264909, 264688, 264556, 264558	264905, 264907, 87168559, 264764	264681, 264685, 264686, 264692	264629	264910, 264686, 264534	263978	264909, 60170394	22278998, 264907, 264681, 264685, 264689, 265020, 264693, 22278000, 22279002, 264566	264906, 264908, 264909, 265006, 264910, 265011, 265017, 264764, 264766, 264767, 264769, 264631, 264634, 264638, 264567, 264486	264488, 264489, 264508, 264509, 264510, 264511, 264512, 264591, 264592, 264601, 264684, 264685, 264769, 264558, 22279002, 264486
	UNGLASSIFIED	UNCLASSIFIED	UNCLASSIFIED			tm7	struct		glycoprotein	kinase	glycoprotein	UNCLASSIFIED
						Contains protein domain (PF00001) - Itm7 7 transmembrane receptor (rhodopsin family)	Contains protein domain (PF00560) - struct Leucine Rich Repeat			Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	
1336 87787890 (2671, 2672) Novel Protein sim. GBank gil465445 sp P33485 VNUA_PRVKA - PROBABLE NUCLEAR ANTIGEN	94312042 (2673, 2674) Novel Protein sim. GBank gil5689471 dbj BAA83019.1 - (AB028990) KIAA1067 protein [Homo sapiens]		80249231 (2677, 2678) Novel Protein sim. GBank gil1176422 (U43194) - rhophilin [Mus musculus]			80089017 (2683, 2684) Novel Protein sim. GBank gi 5019564 emb CAB44507.1 - Contains protein domain (F (AL035542) d.)994E9.5 (hs6M1-17 (novel 7 transmembrane 7 transmembrane receptor receptor (rhodopsin family) (olfactory receptor like) protein)) (rhodopsin family) [Homo sapiens]	80082862 (2685, 2686) Novel Protein sim. GBank gil4557543 ref NP_001384.1 pECM2 - extracellular matrix protein 2		91225546 (2689, 2680) Novel Protein sim. GBank gil2144101 pir 55210 - tricarboxylate carrier - rat (fragment)	1346 80255717 (2691, 2692) Novel Protein sim. GBank gij3881052 emb[CA419523] - (AL023843) predicted using Genefinder; similar to serine/threonine kinase; cDNA EST yk246a12.3 comes from this gene; cDNA EST yk358c10.5 comes from this gene; cDNA EST graph comes from this gene; cDNA EST yk246a12.5 comes	80417393 (2693, 2694) Novel Protein sim. GBank gil4504379 ref NP_003658.1 pHG38 - orphan G protein- coupled receptor HG38	87352335 (2695, 2696) Novel Protein sim. GBank gij3399720 dbj BAA32100 - (AB010999) peptidylarginine deiminase type IV [Rattus novegicus]
		80366114 (2675, 2676)		88316311 (2679, 2680)			80082862 (2685, 2686)	20552559 (2687, 2688)		80255717 (2691, 2692)	80417393 (2693, 2694)	
1336	1337	1338	1339	1340	1341	1342	1343	1344	1345	1346	1347	1348

1349	91225548 (2697, 2698)	1349 91225548 (2697, 2698) Novel Protein sim. GBank gil2144101 pir 155210		UNCLASSIFIED	52646842, 35696286, 22278996, 22278998,
		incarboxylate camer - rat (fragment)	•		22278999, 264259, 29331822, 29331824,
					35696052, 264509, 264905, 264906, 264907.
					264908, 264909, 264511, 265008, 264512,
					264910, 60170831, 264591, 60433438,
					264757, 21906754, 265017, 265018, 264605,
					264760, 264762, 264288, 264766, 264689,
					21906765, 21906766, 21908767, 21906768,
					55811957, 35695917, 265020, 264534,
					264691, 264692, 33657023, 264693,
				ł	33657349, 18108374, 18108376, 35696423,
				**	60170394, 22279000, 22279002, 264563,
1350	87093136 (2699, 2700)			UNCLASSIFIED	52646842, 264259, 29331825, 264908.
			,		264511, 264604, 264288, 21906769, 265020,
					33657182, 33657349, 18108374, 35695855,
					264555, 264558, 18108385, 22279002,
136.	0720 4020, 2004 0200				264486
1001		orserszi (z/ur, z/uz) mover Pfolein sim. GBank gij488/239[gb]AAD32246.1 - (AF064564) BAW protein [Fugu rubnpes]		UNCLASSIFIED	264906, 264907, 264638
1352	80076386 (2703, 2704)			UNCLASSIFIED	264693 263981
1353		95345417 (2705, 2706) Novel Protein sim, GBank oil 2144101 pir (1155710 -			JERORDRE RANJANEO DENONE DENEGO
_		(fricarboxylate carrier - rat (fragment)			SECOND SEASON SEASON SEASON SEASON
					264512 264010 264759 264505 E5911395
					204312, 204910, 204/30, 204330, 33011300, 1
					265011, 264605, 55811150, 264762, 264764,
					264766, 52644229, 56181562, 35695917.
					265022, 33657023, 264693, 35695763,
					60431528, 264629, 263978, 35696423,
					35695855, 264630, 264634, 264635, 264636.
					264637, 264638, 264639, 18108385, 264563,
7367					284564, 264566
<u>.</u>		93330043 (2707, 2706) Nover Frotein Sim. GBank oil4689108inblAAD27763 114E07703 - (AE077030)		UNCLASSIFIED	22278995, 22278999, 29331826, 264906,
		hypothetical 43.2 kDa protein [Homo sapiens]			200000; 3000170£; 21300073; 203011; 87168559 264684 264369 264769 264689
					21906765, 21906768, 52644150, 33657023.
					264692, 264693, 18108374, 83373044.
					87168518, 22279000
1355	88260186 (2709, 2710)	1355 88260186 (2709, 2710) Novel Protein sim. GBank gi 1469199 db BAA09487 -	Contains protein domain (PF00076) - dna_ma_bind	dna_ma_bind	22278996, 22278997, 264259, 66714117,
		(D50928) The KIAA0138 gene product is novel. [Homo	RNA recognition motif. (a.k.a. RRM,	•	264511, 21906754, 265010, 264769, 264689.
		sapiens]	RBD, or RNP domain)		21906765, 21906768, 21906769, 264532,
					27486262, 264629, 264638, 264556, 264638,
					264639, 264482, 264484

1356	95313991 (2711, 2712)	1356 95313991 (2711, 2712) Novel Protein sim. GBank gil 113865 (U40342) - ninein		struct	18108397 22278995 22278996 22278998
		[Mus musculus]			264094, 29331828, 264905, 265008, 265007,
					265008, 265010, 265017, 265018, 265019.
					264764, 18108354, 264689, 21906765,
					265022, 18108364, 35696423, 83373044,
_					18108387
135/	88250258 (2713, 2714)	88260268 (2713, 2714) Novel Protein sim. GBank gij897693jemb CAA90330 -	Contains protein domain (PF01852) -		264259, 29331822, 29331825, 264510,
		(250026) phosphatidylcholine transfer protein [Bos taurus]	START domain		87168559, 265018, 264448, 264288,
					21906765, 21906766, 21906768, 265021.
_					264693, 18108376
1358	38719455 (2715, 2716)	38719455 (2715, 2716) Novel Protein sim. GBank gi[556219 (L36831) - transcription			264757
١		regulator (mus musculus)		1	
339	87771643 (2717, 2718)			UNCLASSIFIED	264907, 264909, 264510, 264511, 264512,
					18108351, 264764, 264534, 33657023,
					18108374, 264634, 264635, 264638, 264639, 18108385, 264488, 264587
1360	87778979 (9710 9790)	87778272 (2719 2720) Movel Droteip eim CBank gil359828210mblC84756121			60422360 264606
	(41,12,12,13)	(V15417) acetateCoA ligase [Coprinus cinereus]		ağıllığa	00452689, 204003
1361	87593527 (2721, 2722)	87593527 (2721, 2722) Novel Protein sim. GBank gij568943 dbj BAA83005.11 -	Contains protein domain (PF00536) - UNCLASSIFIED		35696286, 22278997, 22278999, 264259.
		(AB028976) KIAA1053 protein [Homo sapiens]	SAM domain (Sterile alpha motif)		29331826, 264508, 264509, 264905, 264907,
					264908, 265007, 265009, 33109954,
	•				21906754, 87168474, 265011, 264761,
					264683, 264288, 264766, 264769, 264689,
					21906768, 265020, 265021, 33657023,
					55811576, 35696423, 264634, 60432113.
			-	,	22279002, 264482, 264486
1362	95287961 (2723, 2724)	1362 95287961 (2723, 2724) Novel Protein sim. GBank gij5689411 dbj BAA82989.1 -	Contains protein domain (PF00400) - eph		56182575, 56181686, 60432049, 264259,
		(AB028960) KIAA1037 protein [Homo sapiens]	WD domain. G-beta repeat		29331822, 56182181, 29331827, 35696052
					29331828, 264905, 264906, 264908, 264595.
					55812038, 85658542, 55811150, 264681.
					264288, 264369, 56181562, 60431528,
					55810764, 35696423, 60431850, 264558
1363	85758476 (2725, 2726)	1363 [85756476 (2725, 2726) Novel Protein sim. GBank gi 1130494 (U35776) - ADP-	Contains protein domain (PF01412) - UNCLASSIFIED		264488, 29331826, 264907, 264687, 264689,
		ribosylation factor 1-directed GTPase activating protein	Putative GTP-ase activating protein		264693
_		[Rattus novegicus]	for Arf		
1364	88179488 (2727, 2728)				60432289, 60433356, 60433438, 87168559,
					264603, 18108351, 21906766, 35696423,
					60432113
1365	83003108 (2729, 2730)	83003108 (2729, 2730) Novel Protein sim. GBank gil4589562 dbj BAA76803.1 -		oucogene	264766
		(AB023176) KIAA0959 protein [Homo sapiens]			
1366	87003262 (2731, 2732)	87003262 (2731, 2732) Novel Protein sim. GBank gi[108494 pir] S54495 -	Contains protein domain (PF00153) - transport		265007
		hypothetical protein YPR021c - yeast (Saccharomyces	Mitochondrial carrier proteins		
		Celeviside)			

264488, 52646842, 52646365, 22278995, 56994075, 35696286, 22278996, 22278998, 22278999, 262278999, 262278999, 262278999, 262278999, 264259, 29331824, 66714117, 29331825, 60432289, 35696052, 264905, 264907, 264908, 264909, 264510, 265006, 265007, 265009, 265019, 18108351, 265448, 265017, 265019, 18108351, 265448, 264682, 265019, 18108351, 265020, 265021, 265022, 264534, 60170615, 265002, 265021, 265022, 264534, 60170615, 265002, 265021, 265022, 264534, 60170615, 265002, 265021, 265022, 264534, 264584, 18108376, 33657109, 33657109, 33657109, 33657109, 3265734, 264528, 18108376, 3569423, 35694332, 60170394, 264558, 18108381, 18108385, 56526486, 22279000, 264563,	264567 264259, 29331822, 29331826, 60432289, 29331827, 35696052, 264508, 264905, 264906, 264909, 264510, 265007, 264910, 60433438, 264758, 85658542, 87188559, 284600, 264601, 264760, 264764, 264765, 264768, 52644229, 264689, 35695917, 265020, 265021, 284631, 284632, 264634, 264637, 52644332, 264558, 284639	93373444, 264563, 264566, 264466, 264567 265008, 60432229, 60433356, 33657084, 21906764, 21906769, 264555, 264638, 264559, 264567	22278996, 264259, 29331822, 29331824, 29331826, 264509, 29331827, 35696052, 264508, 264509, 264906, 264906, 264907, 264908, 264909, 264510, 264511, 264512, 264758, 265011, 265019, 264764, 264766, 264768, 21906767, 35695917, 18108362, 35696423, 264638, 264638, 264556, 264638, 264568, 264586, 26	2278996, 264259, 29331822, 29331824, 29331825, 29331825, 29331827, 264905, 29331827, 265908, 265009, 265009, 265009, 265019, 265019, 265019, 264758, 33657084, 85658642, 264448, 3565910, 265010, 265010, 33657109, 33657148, 33657349, 34569854, 33657349, 34569854, 344548, 34569854, 344548, 34569854, 344548, 34569854, 344548, 34569854, 344548,
UNCLASSIFIED		UNCLASSIFIED	potassium_channel	glycoprotein
Contains protein domain (PF01342) - UNCLASSIFIED SAND domain	Contains protein domain (PF00071) - oncogene Ras family			Contains protein domain (PF00335). 4 transmembrane segments integral membrane proteins
i 307 077210 (2733, 2734) Novel Protein sim. GBank gily884088 emb CAB43240.1 - (AL050019) hypothetical protein [Homo sapiens]	94320078 (2735, 2736) Novel Protein sim. GBank gil464561[splP35289[RB15_RAT RAB-15]	86634033 (2737, 2738) Novel Protein sim. GBank gil2062702 (U90550) - butyrophilin [Homo sapiens]	95316910 (2739, 2740) Novel Protein sim. GBank gil5031823 ref NP_005823.1 pKCNM - potassium large conductance calcium-activated channel, subfamily M, beta member 2	95336512 (2741, 2742) Novel Protein sim. GBank gil5032203 ref NP_005714.1 pTSPA - tetraspan 5
0//2/5/ (4/35, 4/34)			85316910 (2739, 2740)	95336512 (2741, 2742)
š	1368	1369	1370	1371

1372	80248517 (2743, 2744)	1372 80248517 (2743, 2744) Novel Protein sim. GBank gil840708 dbj BAA09334 -		collagen	263978
	80499421 (2745, 2746)			UNCLASSIFIED	264769, 21906765, 21906767, 22278999, 264691, 264910, 55812038, 265010, 264681, 264684
1374	95087036 (2747, 2748)	95087036 (2747, 2748) Novel Protein sim. GBank gij111876 pir JC1241 - beta- interferon-induced protein - rat		interferon	264907, 264510, 265007, 264512, 265008, 60432229, 264689, 65274791, 264555, 264557, 83373044, 60432113
		94236942 (2749, 2750) Novel Protein sim. GBank gil5649176[gblpAD03500.2] - (AF051155) G beta-like protein GBL [Rattus norvegicus]	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		52844507, 52845156, 52846842, 52846365, 56182575, 56181686, 22278938, 56994075, 35696286, 22278938, 5294075, 35696286, 22278938, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278939, 22278930, 2245080, 2237826, 224508, 224509, 224509, 224509, 224509, 2244296, 87168474, 87168559, 264500, 265017, 265018, 265019, 25811150, 18108351, 264448, 264764, 26428, 264369, 264429, 21906765, 21906766, 21906767, 21906769, 25811957, 35895917, 265020, 265021, 2264203, 264624, 264634, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264567, 22279000, 22279002, 264563, 264566, 264566, 264566, 264566, 264566, 264567, 22279000, 22279002, 264563, 264566, 264566, 264566, 264566, 264567
1376	87399050 (2751, 275 <u>2)</u>	87399050 (2751, 2752) Novel Protein sim. GBank gil138350lsplp28968IVGLX_HSVEB - GLYCOPROTEIN X PRECURSOR		UNCLASSIFIED	264768, 264769, 35695917, 22278997, 264691, 264259, 29331822, 264693, 35696052, 264508, 264509, 264693, 264605, 264509, 264628, 264900, 264621, 264510, 264511, 264512, 265008, 264630, 264631, 264910, 264632, 264637, 264638, 2646468, 264668, 264688, 264668, 26468
1377	86964242 (2753, 2754)	86964242 (2753, 2754) Novel Protein sim. GBank gil 1663648 (U75321) - chromaffin Contains protein domain (PF00122) - ATPase_associated 29331824, 264591, 265019, 264686, 264768, granule ATPase II homolog [Mus musculus]	Contains protein domain (PF00122) - A	ATPase_associated	29331824, 264591, 265019, 264686, 264768, 55811957, 264693, 22279002

22278994, 22278998, 264093, 264094, 264259, 29331827, 265009, 265018, 265019, 18108351, 264369, 264288, 29148627, 55811957, 264691, 18108368, 33657109, 18108368, 264835, 263981, 18108385	265020				29331824, 29331826, 255007, 50432229, 60433356, 85658542, 265017, 265018,	264685, 264768, 21906766, 35695917,	33657023, 27486261, 27486262, 35695763,	T	Τ		264760, 18108351, 264766, 264769,	35595855, Z64650, Z64656, Z64553, Z64636,	Τ		60432289, 29331828, 264906, 264907,	56182435, 265011, 264681, 60170615, 33657023, 83373044, 264566	D 264591	65274572, 22278999, 264259, 29331826.	29331827, 35696052, 264509, 264907, 264008 264008 264008 264008 2650008 2650008 2650008 2650008 2650008 2650008 2650008 2650008 2650008 2650008 26500008 2650008 265000000000000000000000000000000000000	33657402, 60433438, 264596, 21906754,	87168559, 264600, 265017, 264683,	18108354, 52644229, 21906765, 21906766,	21906767, 21906768, 21906769, 265021,	264558, 60170394, 83373044, 22279000
		nucl_recpt	UNCLASSIFIED	UNCLASSIFIE				Caracon Civil	UNCLASSIFIED				INCI ASSIFIED		tnf		UNCLASSIFIED	glycoprotein						
				Contains protein domain (PF00583) - UNCLASSIFIED	Acetyltransferase (GNAT) family														v			- Address		
1378 87595071 (2755, 2756) Novel Protein sim. GBank gil4107015 dbj BAA36293 - (AB001772) PEM-5 [Ciona savignyi]	85679344 (2757, 2758) Novel Protein sim. GBank gil3252872 (AF035620) - BRCA1- associated protein 2 [Homo sapiens]	87627962 (2759, 2760) Novel Protein sim. GBank gil4837737 gb AAD30662.11- (AF096834) germ cell specific Y-box binding protein [Homo saplens]	88179656 (2761, 2762) Novel Protein sim. GBank gil4731580lgblAAD28508.1 AF12538 - (AF125384) L82A [Drosophila melanogaster]	_									86378788 (2760 2770) Novel Bratein eim GBank oil 2384732 (AE015011) - NAC-1	protein [Rattus norvegicus]	91013049 (2771, 2772) Novel Protein sim. GBank gi 2384910 (AF022982) -	contains similarity to the A-type potassium current class of channel proteins ICaenorhabditis elegans!	87797958 (2773, 2774) Novel Protein sim. GBank gild 160304 jemb[CAA10600] - (A 1132102) HS1 binding protein 3 Mus musculus	95101652 (2775, 2776) Novel Protein sim. GBank	Ψ,	disease resistance protein (Arabidopsis thailana)				
87595071 (2755, 2756)				94847576 (2763, 2764)					8/860398 (2/65, 2/66)				_		91013049 (2771, 2772)		1	+						
1378	1379	1380	1381	1382					1384	<u> </u>			1205	<u> </u>	1386		1387	1388						

1389	91256016 (2777, 2778)	91256016 (2777, 2778) Novel Protein sim. GBank gij5689387 dbj BAA82977.1 -	Contains protein domain (PF00641) - UNCLASSIFIED		65274572, 22278999, 264259, 29331822,
		(AB028948) KIAA1025 protein [Homo sapiens]	Zn-linger in Kan binding protein and others		29146499, 264906, 66712502, 55812038,
					265017, 265018, 265019, 18108351, 264369.
					21906765, 21906766, 21906767, 21906768.
					265020, 265021, 264692, 33657023,
					33657349, 18108370, 18108374, 55811576,
_					264555, 264556, 264557, 60170394,
					83373044, 22279000, 264563, 264564
1307	_	04111916 (2779 2780) Novel Protein sim GBank ail3702295 (AC005783) -		peptidase	52645156, 52646365, 264259, 52645080,
3		R33083 1 (Homo sapiens)			29331825, 29331826, 264906, 52644045,
_					265009, 33657084, 52644296, 87168474,
					87168559, 265017, 265018, 264760, 264682.
				-	264288, 264686, 264687, 56181562,
					52644229, 21906765, 21906769, 35695917.
					265020, 265021, 60170615, 52644150,
					33657023, 27486261, 27486264, 65274791,
					264631, 264555, 52644332, 87168518,
					22279000, 264567
, 36,	10970 1976 (9784 9789)	04507246 (0784 0782) Novel Protein eim GBank	Contains protein domain (PF00709) -		29331826, 29331828, 29331830, 264448,
2	(2012,1012) (2012,216	AIT3469101601P386501P11A1 MOUSE -	Adenylosuccinate synthetase	•	264288, 33657023, 18108365, 264555,
		ADENYLORICONATE SYNTHETASE MUSCLE			264556, 83373044
_		SOCIAL COOCCIENTE STATE TO SECULE			
1302	194311097 (2783 2784)	04311007 (2783 2784) Novel Protein sim GBank gil726286 (U22394) - mSin3A			52646842, 65274572, 22278994, 22278995.
760	23 (5) (5) (5) (5)	Mus musciflies			35696286, 56994075, 22278997, 22278998,
					22278999, 60432049, 264259, 52645080,
					29331822, 29331824, 60432289, 29331826,
					29331827, 35696052, 29331828, 33656970,
	`				264907, 52644045, 265006, 265007, 265008.
					60431735, 60433356, 52646317, 55811386.
					52644296, 265010, 87168559, 265017,
_					264604, 265018, 265019, 264448, 264288,
					264369, 264766, 21906764, 21906767,
					35695917, 265020, 265021, 33657109,
					52645129, 27486261, 27486262, 27486265.
					33657349, 35695763, 18108370, 18108374,
					18108376, 55811576, 35696423, 35695855,
					264636, 52644332, 18108382, 18108385,
					87168518, 60432113, 22279000, 264484,
	_				264566, 18108391
1393	80409472 (2785, 2786)		Contains protein domain (PF00560) - UNCLASSIFIED	UNCLASSIFIED	264763, 264631
			Leucine Rich Repeat		
1394	_			UNCLASSIFIED	264629
136	-	95381471 (2789 2790) Novel Protein sim GBank gil2274845 dbilBAA21534 -		UNCLASSIFIED	265009, 18108381
CACI		Novel Protein sim. Gbain gifz/ +04-5 usilovas 1554 /ORB461) N-WASP [Rathis rathis]			
		(החווםו פחוושנו להעאראו (בפאסרו)			

1396	95363253 (2791, 2792)	1396 95363253 (2791, 2792) Novel Protein sim. GBank gil2135904[pir] 54810 - pHL			22278007 2228000 364250 20324825
		E1F1 - human			£££70331, £££70333, £04£33, £33310£3,
_			_		00432289, 29331828, 29146498, 29146499,
_	-			-	264907, 264908, 29331830, 264909, 265006,
_					265007, 265008, 265009, 60433356, 265010,
					264602, 265017, 265018, 265019, 18108354,
					52644229, 18108358, 21906767, 29148627
					21906768, 21906769, 29148629, 29148784.
					265021, 265022, 18108368, 18108374
	_				58182323 18108385 264563 264567
1397	87631317 (2793, 2794)			UNCLASSIFIED	35696286 264907 66712502 264510
	j				3600017 264607 264602 26606422
1398		91233667 (2795, 2796) Novel Protein sim GRank nil54207389 embl CABAGES 11			33093917, 204092, 204093, 33096423
!					264259, 29331822, 29331824, 29331825,
		Paragraph Protections Protection Protection Protection		`.	29331827, 35696052, 33656970, 87168474,
					265018, 265019, 264682, 264768, 21906767,
					265020, 33657023, 27486261, 55811576,
					264632, 264639, 83373044, 87168518
000,	2020,020,020				22279002
888	87531076 (2797, 2798) 	8/05/10/0 (2/9/, 2/99) Novel Protein sim. GBank gi 2496887 sp Q09232 YQ22 CAEEL - HYPOTHETICAL		UNCLASSIFIED	264768, 18108370, 264555, 264557
		32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III			
400		95419064 (2799, 2800) Novel Protein sim. GBank gil283920 pir S27939 - tensin -	Contains protein domain (PF00017) - UNCLASSIFIED	UNCLASSIFIED	56182575, 22278994, 22278997, 264259.
		chicken	Src homology domain 2		29331822, 29331825, 29331826, 29331827.
					29331828, 264908, 56182435, 264112,
	,				265009, 265011, 265017, 265018, 265019.
					264760, 264762, 264765, 264288, 264685.
					264687, 56181562, 264769, 21906766.
					21906767, 55811957, 264691, 264692,
					264628, 264629, 55811576, 264634, 264555,
					264637, 264557, 264638, 18108381, 264558,
67.	04276270 12004 20021				18108384, 60432113, 22279000
	312203/3 (2001, 2002)	31220373 (2001, 2002) Novel Protein Sim. GBank gij3256185jemb CAA15485j -	Contains protein domain (PF00790) - UNCLASSIFIED	UNCLASSIFIED	65274572, 60432289, 264909, 264758,
440	1000 0000 327	(ALU08635) dJ510H16.1 (Homo sapiens)	VHS domain		264768, 21906769, 22279002
_	93301473 (2003, 2804)	3330 1473 (2003, 2004) Novel Protein sim. GBank gi 1515427 (U57523) - nel	Contains protein domain (PF00008) - tgf	tgf	264905, 264907, 264908, 264909, 264112,
5,		nomolog (Homo sapiens)	EGF-like domain		264693, 33657109, 264634
		94 147933 (2803, 2805) Novel Protein sim. GBank gij5262615 emb CAB45747.1 -			65274572, 66712502, 265017, 264448,
		(ALUSU156) hypothetical protein [Homo sapiens]			264288, 21906765, 21906769, 264693,
740	1404 00035303 (3807 3808)				55811576, 65274791, 60432113
\$	(2002, 1007) cecese			UNCLASSIFIED	65274572, 22278998, 29331822, 29331828,
_					66712502, 265008, 60433438, 265017,
					264693, 18108385

1405	95095068 (2809, 2810)	1405 95095068 (2809, 2810) Novel Protein sim. GBank gil854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]			264488, 56994075, 35696286, 29331822, 29331824, 29331826, 29331828, 35696052, 264508, 264907, 264908, 264510, 264511, 264910, 33657402, 264594, 264768, 264600, 264604, 264762, 18108351, 264764, 33657023, 33657109, 264628, 264634, 22279002, 264563, 264482, 264486, 264567
	87612369 (2811, 2812)	 1406 87612369 (2811, 2812) Novel Protein sim. GBank gil624076 gb AAC96425.1 - (U42580) contains Pro-rich Px motifs: SPKPP (20X), PEPPA (9X); similar to soybean pro-rich cell wall protein, corresponds to Swiss-Prot Accession Number P13993 (Paramecium bursaria Chlorella virus 1) 		collagen	264907, 264605
		Novel Protein sim. GBank gij2827886 (AF015037) - endooligopeptidase A related protein; EOPA related protein [Oryctolagus cuniculus]		UNCLASSIFIED	35696286, 22278999, 264094, 264259, 66714117, 29331826, 29331827, 29331828, 2946498, 264107, 264908, 265006, 265008, 264910, 60433438, 265011, 265017, 18108351, 264448, 264288, 264686, 21906769, 264682, 264686, 264686, 264686, 264686, 264688, 264688, 264688, 264688, 264658, 264511376, 264631, 264557, 266458, 83373044, 18108385, 87168518, 60432113, 22279002
8041		95361477 (2815, 2816) Novel Protein sim. GBank gi 2564953 (AF030001) - unknown [Mus musculus]	Contains protein domain (PF00008) - oncogene EGF-like domain	oncogene	264488, 264489, 35696286, 264109, 264508, 264905, 264906, 264907, 264908, 264907, 264908, 264907, 264908, 264907, 264908, 264909, 265090, 264900, 33657402, 264757, 264758, 265091, 264691, 264687, 264689, 264681, 264691, 264691, 33657023, 264692, 264693, 33657109, 264628, 264629, 36996423, 266455, 264631, 264637, 264638, 56182323, 2645639, 264563, 264563, 264565, 264563, 264564, 264565, 264566, 264566, 264566
1409	66644385 (2817, 2818)	66644385 (2817, 2818) Novel Protein sim. GBank gil2662165 dbj BAA23714 - (AB007902) HH0712 cDNA clone for KIAA0442 has a 574. bp insertion at position 1474 of the sequence of KIAA0442. [Homo sapiens]			264693
1410	86612587 (2819, 2820)	Novel Protein sim. GBank gi[2493790 sp Q60994 ACR3_MOUSE - 30 KD ADIPOCYTE COMPLEMENT-RELATED PROTEIN PRECURSOR (ACRP30) (ADIPOCYTE SPECIFIC PROTEIN ADIPOQ)	Contains protein domain (PF00386) - complement C1q domain	complement	29331826, 264112, 264512, 265009, 265010, 264601, 264686, 264769, 21906767, 263974, 264631, 264566

,,,,					
<u>-</u>	67818641 (2821, 2822)	1411 (6/818641 (2821, 2822) Novel Protein sim. GBank	Contains protein domain (PF00400) - kinasereceptor	kinasereceptor	22278995, 22278997, 22278999, 29331822.
		gij3123135 spjP91343 YM3M_CAEEL - HYPOTHETICAL	WD domain, G-beta repeat		60432289, 29331828, 264907, 265017,
		49.0 KD TRP-ASP REPEATS CONTAINING PROTEIN			265019, 264682, 21906767, 21906768.
		F55F8.5 IN CHROMOSOME I			21906769, 265020, 264690, 264691.
					33657023, 33657109, 27486264, 264628.
1413	94300040 (2022				263972, 264634, 264558, 18108385
	05,400 19 (2023, 2024)			UNCLASSIFIED	264757
<u> </u>	92416559 (2825, 2826)	1413 93416339 (2825, 2826) Novel Protein sim. GBank gi 3879121 emb CAA94370 -	Contains protein domain (PF00023) - homeobox	homeobox	56994075, 29331822, 35696052, 29331828.
		(270310) predicted using Generinder; Similarity to Mouse	Ank repeat		29331830, 264909, 52644045, 264510.
		ankyrin (PIK Acc. No. 537771); cDNA EST EMBL: T01923			52644296, 85658542, 87168474, 265017,
					265018, 264681, 264687, 21906768,
		Thom this gene; conversion this comes from this			35695917, 265020, 52644150, 264692,
_					263967, 27486264, 35695763, 264639,
1414		The state of the s			18108387, 264566
<u> </u>		1940/3000 (2027, 2020) INGVEI PROTEIN SIM. GBANK 91(3252981 (AF068921) - Ras-	Contains protein domain (PF00560) - UNCLASSIFIED		264682, 264683, 265022, 264636
1445	04236048 (2820 2820)	unuming protein SOR-8 (Mus musculus)	Leucine Rich Repeat		
?	94360340 (2023, 4030)	naver respect (2023, 2030) Inover Protein Sim. GBank gij18/1187 (U90439) - unknown			52646365, 56182575, 22278994, 22278995,
		protein (Arabidopsis maliana)			56994075, 22278996, 22278997, 22278998,
					22278999, 264259, 29331822, 29331824,
	-				29331825, 29331826, 29331827, 29331828,
_					29146498, 66712502, 29331830, 52644045,
					264113, 264511, 33657402, 264757,
					21906754, 55811386, 265017, 265018,
_					265019, 264761, 264683, 264369, 264288,
					264686, 264689, 21906766, 21906767,
					29148627, 21906769, 55811957, 265020,
					265021, 264690, 33657023, 65274620,
					52645129, 27486262, 27486264, 60431528,
_					264629, 35695855, 56182323, 264559,
					60432113, 264404, 22279002, 284482

18108392, 18108394, 18108397, 18108398, 22278995, 22278995, 22278996, 22278996, 22278997, 22278998, 22278996, 22278997, 22278998, 22278999, 29331822, 35696052, 29331828, 2946498, 264909, 264113, 265006, 265007, 265008, 265009, 60170831, 264060, 265011, 87168559, 265017, 265018, 265019, 26402, 18108351, 264681, 284763, 264682, 264683, 264683, 264681, 284763, 264689, 21906765, 21906766, 21906767, 29148627, 21906765, 21906767, 29148627, 2644150, 18108361, 264628, 18108370, 264629, 18108362, 264628, 18108370, 264629, 264634, 264636, 56182323, 18108381, 60170394, 18108385, 56526486, 87168518, 22279000	264107, 264448	264259, 264908, 265010, 52644229, 21906764, 21906768, 264690, 264639, 18108388	264259, 60432289, 265006, 87168474, 264288	29331824, 265007, 264563	18108358, 18108396, 18108397, 21906766, 18108398, 21906767, 56182575, 21906768, 21906769, 56182575, 21906768, 21906769, 56181686, 55811957, 35695917, 35695286, 22278996, 22278997, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 264502, 60170615, 29331826, 27486261, 29331822, 3657109, 23357349, 264907, 60431528, 60431250, 264917, 565047, 60431820, 6043229, 60431735, 56182323, 26458, 18108385, 264512, 265017, 265018, 87168559, 60432113, 265017, 265018, 265019, 22279002, 55811150, 264566, 264486, 18108391
struct	ATPase_associated 264107, 264448	UNCLASSIFIED	struct	ATPase_associated	kinase
Contains protein domain (PF00735) - struct					
	87826663 (2833, 2834) Novel Protein sim. GBank gil4958935 dbj BAA78095.1 - (AB027570) suppressor of potassium transport defect 3 [Rattus norvegicus]		87757168 (2837, 2838) Novel Protein sim. GBank gi 2072294 (U95097) - mitotic phosphoprotein 43 [Xenopus laevis]	Novel Protein sim. GBank gi[5174421[ref]NP_066023.1[pCPNE - copine VI (neuronal)	94746986 (2841, 2842) Novel Protein sim. GBank gij3876090jemb CAA93459.11- (Z69635) Similarity to Yeast uridine kinase (SW:URK1_YEAST); cDNA EST EMBL:Z14695 comes from this gene; cDNA EST CEMSE17F comes from this gene; cDNA EST EMBL:D67355 comes from this gene; cDNA EST yk209h1.5 comes from this gen
			$\neg \neg$		
<u> </u>	1417	1418	1419	1420	1421

	881/8/// (2843, 2844)	081 / 08 / 1 (2843, 2844) Novel Protein Sim. GBank gi 4505939 ref NP_000928.1 pPOLR - polymerase (RNA) II (DNA directed) polypeptide A (220kD)		rnapolymerase	35811576, 264555, 264557, 87168518
1423	86997762 (2845, 2846)			UNCLASSIFIED	264686, 264489, 264692, 264594, 264603, 265018, 264908
1424	95201610 (2847, 2848)	95201610 (2847, 2848) Novel Protein sim. GBank gil437181 (U02289) - GTPase-activating protein [Caenorhabditis elegans]	Contains protein domain (PF00620) - struct RhoGAP domain	struct	29331622, 29331625, 29331627, 29146498, 264905, 264906, 264908, 264909, 265007, 264910, 265009, 33109954, 265010, 87168559, 265019, 264766, 264687, 21906765, 21906767, 21906767, 21906767, 21906761, 264691, 264692, 56526486, 22279002, 264563
1425				UNCLASSIFIED	265007, 264558
1426	94322115 (2851, 2852)	94322115 (2851, 2852) Novel Protein sim. GBank gi 2078441 (U56964) - weak similarity to S. cerevisiae intracellular protein transport protein US)1 (SP:P25386) [Caenorhabditis elegans]		UNCLASSIFIED	264488, 60424179, 35696286, 22278997, 22278998, 22278999, 264259, 60432049, 29331822, 29331825, 56182435, 264910, 60433356, 60433438, 21906756, 21906767, 21906767, 21906769, 265022, 23657109, 18108376, 26526486, 22279002, 264482
1427	91227510 (2853, 2854)	1427 91227510 (2853, 2854) Novel Protein sim. GBank gi 5616074 gb AAD45616.1 AF06194 - (AF061943) protate- derived STE20-like kinase PSK [Homo sapiens]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	22278994, 56994075, 22278997, 29331828, 29331830, 264828, 265006, 265007, 265008, 265009, 264083, 264288, 18108354, 21906768, 29148629, 33657023, 18108374, 35695855, 83373044, 22279002, 264564
1428	94323008 (2855, 2856)	94323008 (2855, 2856) Novel Protein sim. GBank gif138350 sp P28968 VGLX_HSVEB - GLYCOPROTEIN X PRECURSOR		glycoprotein	56181686, 264259, 264907, 265007, 265009, 264595, 265010, 264686, 65274620, 264629, 65274791, 22279002, 264566
1429	87888689 (2857, 2858)			UNCLASSIFIED	264112, 264595, 265017, 265019, 21906765. 263977, 264555
1430		94735021 (2859, 2860) Novel Protein sim. GBank gil1181619 dbj BAA11565 - (D82364) a variant of TSC-22 [Gallus gallus]			264094, 29331824, 264591, 264593, 265018, 264681, 21906765, 21906767, 65274620, 65811576, 264639, 87168518, 22279002
1431		80429081 (2861, 2862) Novel Protein sim. GBank gij5420389jemb CAB46680.1 - (AJ243460) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	264768, 265020, 264691, 264693, 264638
1432		87463004 (2863, 2864) Novel Protein sim. GBank gil414797 (L18966) - pyruvate dehydrogenase phosphatase [Bos taurus]		phosphatase	18108394, 29146498, 265007, 60433438, 264763, 29148629, 263969
1433		97605403 (2865, 2866) Novel Protein sim. GBank gi[2460316 (AF022147) - uterus- ovary specific putative transmembrane protein [Rattus norvegicus]	Contains protein domain (PF00100) - UNCLASSIFIED Zona pellucida-like domain	UNCLASSIFIED	264259, 264510, 264591, 264603, 264565
1434	85713730 (2867, 2868)			UNCLASSIFIED	264682, 264691

22278997, 22278998, 22278999, 60432049, 264259, 29331822, 29331824, 66714117, 29331826, 33656970, 264508, 264905, 66712502, 29331830, 264909, 265007, 265008, 264910, 265009, 60433356, 60433438, 264596, 21906754, 265010, 265017, 265018, 265019, 18108351, 264762, 265017, 265018, 265019, 18108351, 264628, 1810876, 265020, 264628, 18108374, 18108374, 524559, 83373044, 87168518, 60432113, 22279000, 22279002, 374563, 264648, 264648, 264648, 264659,	263978, 284557, 264559	22278997, 66714117, 29331826, 264907, 56182435, 265009, 18108351, 264692, 264693	264488, 264508, 264906, 264909, 264757, 284600, 264601, 264605, 264768, 264769, 264690, 35688423, 264588, 264565	264489, 18108394, 65274572, 56182575, 22278994, 22278996, 35696286, 22278997, 22278998, 22278999, 264259, 60432049, 29331822, 29331824, 60432289, 29331826, 29331827, 29331824, 60432289, 29331826, 29331837, 29331828, 35686052, 33556970, 264107, 264508, 264507, 66712502, 29331830, 56182435, 264511, 265006, 265007, 265009, 60432229, 60433438, 264595, 55612038, 55811386, 265011, 265017, 265019, 264019, 18108354, 26448, 18108354, 264187, 265020, 265021, 265022, 33657109, 18108370, 18108374, 55810764, 35695855, 264634, 264638, 56182323, 83373044, 18108387, 81168518, 60432113, 22279000, 26488	264887, 264259, 264906, 264907, 264908, 264909, 265008, 265010, 265017, 265018, 265019, 18108351, 264369, 265020, 33657023, 33657109, 60431528, 55811576, 264635
Iranscriptfactor	UNCLASSIFIED	UNCLASSIFIED	kinase	cathepsin	UNCLASSIFIED
			Contains protein domain (PF00595) - PDZ domain (Also known as DHR or GLGF).	Serine carboxypeptidase	
1435 94708213 (2869, 2870) Novel Protein sim. GBank gij3970850 dbj BAA34789.1 - (AB015330) HRIHFB2007 [Homo sapiens]	86635024 (2871, 2872) Novel Protein sim. GBank gij3183977 jemb CAA39515 - (X56041) protein Htf9C (Mus musculus)	87631082 (2873, 2874) Novel Protein sim. GBank gi 2496887 sp Q09232 YQ22_CAEEL - HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III	83344280 (2875, 2876) Novel Protein sim. GBank gi[1805906 (AD000092) - hypothetical human serine-threonine protein kinase R31240_1 [Homo sapiens]	91231894 (2877, 2878) Novel Protein sim. GBank gij3876299 emb CAA94892 - (Z71180) similar to BPTI/KUNIT2 inhibitor domain; cDNA EST EMBL:D68293 comes from this gene; cDNA EST yk448h4,5 comes from this gene; cDNA EST yk448h4,3 comes from this gene; cDNA EST yk448h4,3 comes from this gene [Caenorhabdi	(AB007902) HH0712 cDNA done for KIAA0442 has a 574-bp insertion at position 1474 of the sequence of KIAA0442. [Homo sapiens]
94708213 (2869, 2870)	- 1	87631082 (2873, 2874)		91231894 (2877, 2878)	
4	1436	1437	55	4.50 0.54 0.54	

18108392, 264488, 263994, 264489, 5689255, 25278994, 22278994, 22278995, 56994075, 35696286, 22278994, 22278999, 60432049, 264259, 29331822, 29331824, 56182181, 66714117, 29331822, 29331824, 56182181, 66714117, 29331826, 264508, 264905, 264509, 264909, 264509, 264909, 264509, 264909, 264509, 264909, 264509, 264909, 264593, 39657402, 60433438, 264592, 264691, 264592, 264691, 264592, 264691, 264602, 265017, 264604, 265018, 264609, 265019, 264604, 265018, 264609, 265019, 264604, 265018, 264609, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 264009, 35695763, 264009, 35695763, 264031, 264634, 264034, 264639, 264639, 264639, 264631, 264634, 264539, 264539, 264539, 264539, 264539, 264569, 264531, 264639, 264564, 26566, 264569, 264569, 264569, 264563, 264569, 264569, 264563, 264569, 26	264906, 265007, 264693, 264558	52645080, 264691, 264628, 264555	264259, 29331822, 29331824, 66714117, 29331828, 264288, 35695917, 33657023, 264635, 60170394	264369	264369	18108396, 35696286, 22278997, 66714117, 29331828, 265009, 264758, 265018, 264288, 21906766, 21906767, 264692, 264634,	29331827, 264509, 264909, 265008, 264595, 18108357, 18108385, 264586
• helicase	transcriptfactor	struct	UNCLASSIFIED	UNCLASSIFIED			
Contains protein domain (PF00646) - helicase F-box domain.		Contains protein domain (PF00435) - struct Spectrin repeat				Contains protein domain (PF01846) - FF domain	3980411 (AC004561) - putative Contains protein domain (PF00439) - sis thaliana]
	83367491 (2883, 2884) Novel Protein sim. GBank gij5103027[dbj BAA78765.1 - (AB023419) mSox7 [Mus musculus]		O/O2U4/8 (2881, 2888) Novel Protein sim. GBank gij3874447[emb]CAB02772] - [Z81039) predicted using Genefinder; cDNA EST EMBL:T01209 comes from this gene; cDNA EST yk278a11.5 comes from this gene; cDNA EST yk278a11.5 comes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.5 com.	Novel Protein sim. GBank gil2959886jemb CAA11022 . [AJ222968] L-perjaxin (Mus musculus)	85079268 (2891, 2892)	1/AF13544 - (AF135440) [Mus musculus]	94990477 (2895, 2896) Novel Protein sim. GBank gij3980411 (AC004561) - putative (protine-rich protein [Arabidopsis thaliana]
-1 95317662 (2881, 288 <u>2</u>		8 87109935 (2885, 2886)		$\overline{}$		0094303K (K083, K084)	94990477 (2895, 2896)
-	442	443	4	1445	1446		448

1449	87860859 (2897, 2898)			UNCLASSIFIED	66714117, 264908, 264908, 264591, 264601, 264764
1450	87458696 (2899, 2900)			UNCLASSIFIED	35696286, 35696052, 265008, 265009, 60170831, 33109954, 264683, 264689, 35696423, 35695855, 56526486
1451	87797970 (2901, 2902)	87797970 (2901, 2902) Novel Protein sim. GBank gil4160304 emb CAA10600 - (AJ132192) HS1 binding protein 3 [Mus musculus]		UNCLASSIFIED	29331826, 264683, 264693, 263978, 264630
1452	_	85692899 (2903, 2904) Novel Protein sim. GBank gi 2832906 dbj BAA24608 1 - (D89340) dipeptidyl peptidase III [Rattus norvegicus]		peptidase	264681, 33657023, 264629
1453		86130434 (2905, 2906) Novel Protein sim. GBank gi 728831 sp P39188 AU1_HUMAN - !!!! ALU SUBFAMILY J WARNING ENTRY !!!!		kinase	264510, 264768
1454	11204696 (2907, 2908)	-			264556
1455				UNCLASSIFIED	29331822, 66714117, 29331825, 264905, 29331830, 265006, 265008, 265009, 265011, 265019, 18108351, 21908768, 33857109, 18108376, 264632, 56182323, 87168518
1456		86320218 (2911, 2912) Novel Protein sim. GBank gil729230 sp P41004 CUT3_SCHPO - CHROMOSOME SEGREGATION PROTEIN CUT3		transport	22278995, 22278996, 22278997, 22278998. 22278999, 29331827, 264107, 265017, 21906765, 21906766, 21906767, 21906769. 29148629, 18108370, 22279000
1457				UNCLASSIFIED	264107, 264566
1458		87800460 (2915, 2916) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73.		transport	56182575 22278999 60432049 264259
					29331826, 25331827, 29331828, 264102, 264102, 264101, 265009, 60432229, 265019
					200420, 200372, 200310, 204000, 22273002. 204566
1459		95360920 (2917, 2918) Novel Protein sim. GBank gij5524687 gblAAD44333.1 AF15935 - (AF159356) Munc13-C2 domain 4 protein [Rattus norvegicus]	Contains protein domain (PF00168) - kinase C2 domain	kinase	22278997, 264259, 29331824, 29331826, 29331827, 29331828, 265017, 265018, 264760, 264682, 26448, 264288, 264766, 265021, 264692, 39557023, 33657109, 3669585
1460	95354602 (2919, 2920)			UNCLASSIFIED	29331822, 264591, 55811957, 264691, 264693, 65274620
1461	94741513 (2921, 2922)	94741513 (2921, 2922) Novel Protein sim. GBank gi 1707274 (U80931) - strong similarity to class-III of ovridoxal-phoshate-dependent	Contains protein domain (PF00202) - gaba Aminotransferases class-III pyridoxal	gaba	22278997, 29331822, 35696052, 265009, 264758, 265017, 265018, 265019, 264760
		aminotransferases [Caenorhabditis elegans]	phosphate		264369, 264687, 21906765, 21906768, 265022, 33657109, 27486261, 264555, 83373044
1462	87732018 (2923, 2924)			UNCLASSIFIED	264555, 264556
1463	88090605 (2925, 2926)	1463 88090605 (2925, 2926) Novel Protein sim. GBank gil1770466 emb CAA66912 -	Contains protein domain (PF00385) - struct	struct	60432049, 264259, 29146499, 264906,
		(X98259) M-phase phosphoprotein 8 [Homo sapiens]	Chromo' (CHRromatin Organization		264907, 264512, 265017, 264763, 264766,
			MOdifier) domain		18108370, 18108374, 264636, 18108385, 18108388

UNCLASSIFIED 264569, 22278995, 22278996, 22278997, 22278998, 29331824, 29331825, 29331822, 29331824, 29331825, 264605, 20278909, 265007, 265007, 33657402, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906769, 35696423, 35695855, 264630, 264630, 264631, 264639, 264630, 2	glucoamylase 264488, 22278994, 56994075, 60432049, 264259, 56182181, 60432289, 29331827, 25644045, 264511, 265007, 265008, 264596, 55812038, 55811386, 264600, 264602, 265017, 265018, 264604, 265019, 18108351, 18108354, 56181562, 21906769, 265021, 33657023, 33657182, 55811576, 264557, 18108382, 60432113	UNCLASSIFIED 264512, 265017, 264689, 264558	struct 264683, 264636	22278998, 264758, 265018, 265019, 21906769, 265020, 33657109, 22279002		UNCLASSIFIED 18108394, 18108398, 56182575, 264259, 29331824, 29331824, 29331825, 60432289, 264907, 264909, 265007, 264910, 265009, 264907, 264909, 265007, 264910, 265009, 264910, 264911957, 265011, 18108361, 21906769, 55811957, 264637, 264557, 18108381, 56182323, 18108382, 83373044, 18108384, 18108388, 87168518, 60432113, 264404, 22279002, 264487, 264567, 264487	10153) - transport
1464 87620482 (2927, 2928) Novel Protein sim. GBank gij3874447 emb CAB02772	87425192 (2929, 2930) Novel Protein sim. GBank gil4589598 dbj BA476821.1 - (AB023194) KIAA0977 protein [Homo sapiens]		87614328 (2933, 2934) Novel Protein sim. GBank gi[2246532 (U93872) - ORF 73, contains large complex repeat CR 73 [Kaposi's sarcoma- associated herpesvirus]	Novel Protein sim. GBank gij4507241 ref NP_003137.1 pSSRP - structure specific recognition protein 1	79236174 (2937, 2938) Novel Protein sim. GBank gil 1906596 (U81788) - kinesin-73 [Drosophila melanogaster]	Novel Protein sim. GBank gi 5649170 gb AAD43131.2 AF15909 - (AF159092) syld709613 protein [Homo sapiens]	87826842 (2941, 2942) Novel Protein sim. GBank gij3876146[emb]CAB01750] - Contains protein domain (PFG (278542) similar to Mitochondrial carrier proteins: cDNA EST EMBL:T01651 comes from this gene [Caenorhabditis]
1464 87620482 (2927, 2928) NG (2 E) (2 E) (3 E) (4 E) (4 E) (5 E) (5 E) (5 E) (6 E) (7 E)		-		1468 95342862 (2935, 2936) No gil	1469 79236174 (2937, 2938) No		1471 87826842 (2941, 2942) No (Z.

1472		1472 87756616 (2943, 2944) Novel Protein sim. GBank gl 4680707 gb AAD27743.1 AF13296 - (AF132968) CGI-34 protein [Homo sapiens]		UNCLASSIFIED	264905, 18108351, 21906765, 264486
1473	Ī	87791609 (2945, 2946) Novel Protein sim. GBank gil3688780 (AF042180) - testis-	Contains protein domain (PF00956) - IMHC	MHC	18108394, 22278995, 56994075, 22278999.
		specific Y-encoded-like protein (Mus musculus)	Nucleosome assembly protein (NAP)		29331822, 29331824, 66714117, 29331825,
					29331826, 35696052, 264906, 264907,
					56182435, 265007, 264758, 265018, 265019,
					264760, 264764, 264288, 264685, 264686,
					264768, 21906769, 55811957, 265021,
					264691, 264693, 264629, 55811576, 264634,
					264638, 56182323, 22279002, 264566,
					264486
1474	85800989 (2947, 2948)	85800989 (2947, 2948) Novel Protein sim. GBank	Contains protein domain (PF00400) - struct	struct	264488, 35695917, 35696286, 264692,
		gij2494890jspjQ92176jCORO_BOVIN - CORONIN-LIKE	WD domain, G-beta repeat		33657023, 264693, 33657109, 35696052,
		PROTEIN P57			264508, 264905, 264906, 264907, 264629,
					264908, 264909, 35696423, 35695855,
					264511, 264910, 264632, 264634, 264635,
					264636, 264637, 264556, 264557, 264639,
					264758, 60432113, 264604, 264605, 264565,
					264566, 264764, 264486, 264685, 264766
1475	86871935 (2949, 2950)		Contains protein domain (PF00041) - UNCLASSIFIED	UNCLASSIFIED	264681, 264682, 264288, 264566
			Fibronectin type III domain		
1476	87548855 (2951, 2952)	1476 87548655 (2951, 2952) Novel Protein sim. GBank	Contains protein domain (PF00147) - glycoprotein	glycoprotein	60424179, 56181686, 29331824, 60424269,
		gi 4757752 ref NP_004664.1 pANGP - angiopoietin 3	Fibrinogen beta and gamma chains,		29331826, 35696052, 264508, 264905,
_			C-terminal globular domain		264906, 264907, 264908, 264909, 264512,
					265007, 265008, 265009, 264910, 33657402,
					264595, 264596, 55812038, 265011, 264601,
					264762, 18108351, 264288, 264369, 264685,
					264766, 264689, 55811957, 264691, 264692,
					264693, 18108370, 60431528, 18108374,
					35696423, 264634, 264635, 264636,
					60431850, 264555, 264638, 264557, 264639,
					18108382, 18108388, 60432113, 22279002
1477		87774279 (2953, 2954) Novel Protein sim. GBank		UNCLASSIFIED	264259, 264107, 264905, 265008, 265010,
		gil2498308 sp Q60870 DP1_MOUSE - POLYPOSIS LOCUS			265011, 264682, 264288, 265020, 265021,
		PROTEIN 1 HOMOLOG (182 PROTEIN HOMOLOG) (GP106)			263974
1478	1478 11754412 (2955, 2956)				264686

1479 [91640140 (2957, 2958) Novel Protein sim. GBank	Novel Pro		d	peptidase	56182575, 22278995, 22278996, 22278998.
gij5499741jgbjAAD43978.1jAF15296 - (AF152961)	gi[5499741]gb AAD43978.1 AF1	F15296 - (AF152961)			22278999, 29331822, 29331824, 66714117, 264906, 264907, 56182435, 265006.
kDa subunit [Homo sapiens]	kDa subunit [Homo sapiens]				60170831, 33657402, 264758, 33108954,
					21906/54, 2650117, 265013, 264448, 264268, 264767, 264687, 52644229, 21906764,
					264689, 21906765, 21906768, 21906769, 266030, 266031, 60170615, 264691
					33657023, 33657109, 33657182, 27486261.
					27486262, 33657349, 18108370, 60431528.
		_			263976, 55811576, 264556, 264557,
					001/0394, 8/100310, 204404, 222/3000, 22279002, 264563, 264482
1480 94312412 (2959, 2960) Novel Protein sim. GBank gi 3550		3550456jemb CAA06329.1 -		UNCLASSIFIED	18108394, 65274572, 56182575, 22278995.
(AJ005073) Alix [Mus musculus]	(AJ005073) Alix [Mus musculus]				33090260, 30934073, 22778990, 22778998, 22278999, 264091, 264259.
					35696052, 29146499, 264103, 264105,
-					264108, 264907, 52644045, 264112, 265007,
					263008, 263008, 60433339, 60433439. 264598, 33109954, 33657084, 52644296,
					87168474, 265010, 87168559, 265017,
				-,-	265018, 265019, 264448, 264682, 264683,
					264769, 21906765, 21906766, 21906767,
					21906768, 21906769, 265020, 265021, 60170614, 52644150, 33657109, 33657182,
					263972, 35695855, 264557, 263981,
		•			83373044, 18108385, 87168518, 264566, 264089, 264486
92000 0000 00000 100000 10000 0000000000	Manal Designation of the Company			MHC	265006, 265007, 265010, 18108374
₹	₹	.F14679 - (AF146793) PFT27			
1482 85320442 (2963, 2964) Novel Protein sim. GBank gi[4585372]gb[AAD25403.1]AF1 Inhibitory factor-1 Mus musculu	Novel Protein sim. GBank gi 4585372 gb AAD25403.1 AF1 inhibitor factor-1 Mus musculu	:92 - (AF122923) Wnt	Contains protein domain (PF00008) - UNCLASSIFIED EGF-like domain	UNCLASSIFIED	264908, 264910, 264758
1483 94115503 (2965, 2966) Novel Protein sim. GBank gil5354	Novel Protein sim. GBank gi[535	428 (U13736) - calmodulin-	535428 (U13736) - calmodulin- Contains protein domain (PF00036) - struct	struct	264259, 29331822, 52645080, 29331825,
like protein [Pisum sativum]	like protein [Pisum sativum]		EF hand		29331826, 33656970, 29331830, 265007, 55812038, 33109954, 265017, 264288,
					21906768, 21906769, 264636, 18108380, 87168518, 22279000
1484 94131544 (2967, 2968) Novel Protein sim. GBank gi 191	Novel Protein sim. GBank gi[191	1911774 bbs 180090 -		UNCLASSIFIED	264489, 35696286, 264259, 264107, 264909,
(S83364) putative Rab5-Interacting protein (clone L1-57) [himan Hel a cells. Peotide Partial, 122 aal [Homo	(S83364) putative Rab5-interacti	acting protein (clone L1-57) Partial, 122 aal [Homo			265008, 60433356, 3365/402, 50433436, 264288, 21906765, 21906766, 29148627,
					33657023, 27486262, 18108374, 35696423, 83373044, 60432113
1485 Roto4441 (2069 2070) Novel Protein cim GBank	Novel Protein sim GBank		Contains protein domain (PF00225) - struct	struct	264369, 265020, 18108374
gi[35602029]gbb/2803.1 AF15511 - (AF155117) NY-REN Kinesin motor domain	gi[5360129]gb AAD42883.1 AF1	15511 - (AF155117) NY-REN	Kinesin motor domain		
loz dinigeri (nomo saprens)	loc amigen (nomo sapiens)				

1488	041250EE /2071 2072	Marrol Destruit Charles all actions of the angelone			
<u> </u>	34123000 (23/1, 23/2)	1400 STILLOOU (LSTI), 23/2/ NOVEL FIDERIN SHIT, GBARK BIJ4369310 DAILY OF BUILL	Contains protein domain (PF00059) - kinase	kinase	56182575, 22278999, 264906, 264907,
		(ABUZ3133) KIAAU936 protein [Homo sapiens]	Eukaryotic protein kinase domain		21906754, 87168474, 265017, 265019,
					18108351, 264288, 265020, 264566
1487	86452711 (2973, 2974)	1487 86452711 (2973, 2974) Novel Protein sim. GBank gij5019275 emb CAB4431.1 -		synthase	21906754, 264486
		(AJ132751) xenoblotic/medium-chain fatty acid:CoA ligase			
		form XL-III [Bos taurus]			
1488	87732026 (2975, 2976)	87732026 (2975, 2976) Novel Protein sim. GBank	Contains protein domain (PF01443) - fgf	fat	264686, 264769, 264689, 264692, 264693
		gi[5712131]gb[AAD47379.1[AF12049 - (AF120499) DEM1	Viral (Superfamily 1) RNA helicase	1	264509, 264906, 264907, 18108370, 264908
		protein [Homo sapiens]			264629, 264909, 264510, 265006, 264512.
					265007, 265008, 265009, 264555, 264556,
					264557, 264558, 264762, 264564, 264682
1469	95104277 (2877, 2978)	95104277 (2977, 2978) Novel Protein sim. GBank	Contains protein domain (PF00047) - prostaglandin	prostaglandin	21906767, 22278999, 265022, 264259,
		gij2497303jspjQ62786jFPRP_RAT - PROSTAGLANDIN F2-/Immunoglobulin domain	Immunoglobulin domain		264693, 29331824, 29331825, 29331826,
		ALPHA RECEPTOR REGULATORY PROTEIN			29331827, 29331828, 264103, 263972,
		PRECURSOR (PROSTAGLANDIN F2-ALPHA RECEPTOR			66712502, 35696423, 35695855, 265007,
		ASSOCIATED PROTEIN)			265008, 265009, 83373044, 21906754,
					56526486, 265017, 264563, 18108351,
[264564, 264566, 264369, 264288
1490	1490 87390127 (2979, 2980)			UNCLASSIFIED	56182575, 264259, 29331822, 29331824,
					66714117, 29331827, 29331828, 264508,
					264905, 66712502, 265007, 265008, 264594,
					33657402, 55812038, 87168474, 265018,
_					18108351, 264369, 264288, 264769, 264689,
					21906767, 21906768, 55811957, 60170615,
·					33657109, 35695855, 264635, 60170394,
					56526486, 22279002, 264563
1491	83594305 (2981, 2982)	83594305 (2981, 2982) Novel Protein sim. GBank gi[295671 (L11275) - selected as		UNCLASSIFIED	265007, 264448, 18108372, 264558.
		a weak suppressor of a mutant of the subunit AC40 of DNA			56182323
		dependant RNA polymerase I and III (Saccharomyces			
		(cerevisiae)			
1492	85805363 (2983, 2984)	85805363 (2983, 2984) Novel Protein sim. GBank gij1656005 (U71205) - rit Mus	Contains protein domain (PF00071) - oncogene		22278997, 22278998, 29331822, 264907
		musculus	Ras family		66712502

264488, 52646365, 65274572, 56182575, 22278994, 35696286, 56994075, 22278999, 60432049, 29331824, 29331828, 35696652, 264508, 284906, 52644045, 264909, 56170831, 33657402, 55812038, 265009, 265011, 265017, 265018, 265019, 55811150, 26448, 264682, 264685, 264686, 52644229, 21906765, 21906766, 21906769, 21906765, 21906765, 21906769, 265020, 265021, 60170615, 52644150, 33657102, 31657349, 27486261, 27486262, 27486264, 33657349, 27486262, 32695763, 18108370, 264629, 18108374, 52844332, 56182323, 87168518, 22279002, 264564, 264566, 264567	264907, 265009, 264769, 18108370. 55811576, 264639, 264565, 264486	22278997, 264259, 264906, 264907, 265009, 264594, 33657084, 265017, 264760, 264448, 33657109, 264634, 56526486, 264563, 264565, 264566, 264486, 264567, 2645767, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264577, 26477, 264777, 264777, 264777, 264777, 264777, 264777, 264777, 264777, 264777, 264777, 264777	35696286, 264906, 265019, 264693	264910, 264758, 265011, 264764, 264288, 264690, 264634, 264635, 56526486	264488, 65274572, 29331822, 265017, 265018, 21906765, 29148627, 29148629, 18108374, 264637, 264638, 264567	263978, 264566	22278999, 264769, 18108379	264559	264508, 264112, 264604, 264684, 52644150. 55811576, 264632, 264556, 264638, 56182323, 264563, 264486	29331822, 265007, 264369
UNCLASSIFIED	kinase	transcriptfactor	UNCLASSIFIED	kinase	oncogene	glucoamylase	UNCLASSIFIED	UNCLASSIFIED		- transferase
	Contains protein domain (PF01352) - kinase KRAB box	Contains protein domain (PF01352) - transcriptfactor KRAB box		Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain						Contains protein domain (PF00535) - transferase Glycosyl transferases
1493 91677215 (2985, 2986) Novel Prolein sim. GBank gij5689515 dbj BAA83041.1 - (AB029012) KIAA1089 protein [Homo sapiens]	<u>}</u>	87605267 (2989, 2990) Novel Protein sim. GBank gil4589588 db BAA76816.1 (AB023189) KIAA0972 protein [Homo sapiens]	87784322 (2991, 2992) Novel Protein sim. GBank gil5420387 emb CAB46679.1 - A_1243459 proteophosohoolycan ILeishmania major	81695428 (2993, 2994) Novel Protein sim. GBank gij3874925jembjCAA92291 - (Z68296) Similarity to Mouse A-RAF proto-oncogene serine/threonine-protein kinase (SW:KRAA_MOUSE); cDNA EST EMBL:D27610 comes from this gene; cDNA EST EMBL:T01018 comes from this gene; cDNA EST FMBL.T033256 comes from this gene; cDNA EST		86451589 (2997, 2998) Novel Protein sim. GBank gil2570198 (U54556) - microfilarial sheath protein SHP3 [Litomosoides sigmodonlis]	-	85795297 (3001, 3002) Novel Protein sim. GBank gil2078483 (U43200) - antifreeze glycopeptide AFGP potyprotein precursor (Boreogadus saida)	_	87012701 (3005, 3006) Novel Protein sim. GBank gil3900855 (AC004874) - similar to N-acetylgalactosaminyltransferase; similar to Q07537 (PID:g1171989) [Homo sapiens]
91677215 (2985, 2986) (87605265 (2987, 2988)	87605267 (2989, 2990)	87784322 (2991, 2992)	81695428 (2993, 2994)	90934938 (2995, 2996)		R0499386 (2999 3000)		80206141 (3003, 3004)	87012701 (3005, 3006)
1493	1494	1495	1496	1497	1498	1499	1500	505	1502	1503

1504	1504 79640051 (3007, 3008)		Contains protein domain (PF00023) - UNCLASSIFIED	UNCLASSIFIED	264693
1505	86102672 (3009, 3010)	1505 86102672 (3009, 3010) Novei Protein sim. GBank gil4753775 emb CAB41970.1 - (AJ132545) protein kinase [Homo sapiens]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	29331826, 35696052, 264509, 264906, 264907, 264908, 264909, 264511, 264910, 25812038, 264759, 264763, 264764, 264689, 285695917, 265022, 33657109, 18108374, 286331, 264634, 26464, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 26664, 266
1506		94143219 (3011, 3012) Novel Protein sim. GBank gi[1304201 dbj BAA06170] - (D29766) atternatively spliced product [Rattus norvegicus]	Contains protein domain (PF00018) - glycoprotein SH3 domain	glycoprotein	5574572, 56182575, 56994075, 22278997, 22278998, 264091, 264099, 264099, 264099, 264099, 264099, 264099, 264099, 264099, 264099, 264090, 26431820, 29331820, 264106, 29331830, 264908, 56182435, 264110, 264511, 264512, 265017, 265018, 264681, 18108354, 264369, 21906754, 87168559, 264600, 265017, 266018, 264681, 18108354, 2643687, 264687, 21906768, 21906768, 29148629, 52644150, 33657023, 4108376, 652747991, 56182323, 244558, 264588, 2645
1507	83738250 (3013, 3014)	83738250 (3013, 3014) Novel Protein sim. GBank gil5689513 db BAA83040.1		helicase	60432113, 22279000, 264565 264639
1508	11618758 (3015, 3016)	(Abotatori) NAM 1000 protein (Homo sapiens)			264503
1509	87318451 (3017, 3018)	87318451 (3017, 3018) Novel Protein sim. GBank gil5031975 ref NP_005875.1 pPAK4 - protein kinase related Eukaryotic protein kinase domain to S. cerevislae STE20, effector for Cdc42Hs	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264259, 29331822, 29331824, 29331828, 264591, 33109954, 264563
1510	95362643 (3019, 3020)	95362643 (3019, 3020) Novel Protein sim. GBank gi 113161 sp P28614 ACOR_ALCEU - ACETOIN CATABOLISM REGULATORY PROTEIN		UNCLASSIFIED	264259, 29331822, 265007, 18108374, 264556
1511		88318073 (3021, 3022) Novel Protein sim. GBank gij728831 sp P39188 ALU1_HUMAN - !!!! ALU SUBFAMILY J WARNING ENTRY !!!!		UNCLASSIFIED	264259, 29331822, 29331824, 29331827, 264509, 284907, 264510, 264511, 265007, 264512, 265008, 87168559, 264288, 265022, 33657023, 35695855, 264637, 264638, 24563
1512		95345390 (3023, 3024) Novel Protein sim. GBank gil4559353[gb]AAD23014 1 AC00658 - (AC006585) putative RIO1/ZK632.3/MJ0444 family extragenic suppressor protein [Arabidopsis thaliana]	Contains protein domain (PF01163) - RIO1/ZK632.3/MJ0444 family		52645156, 18108396, 56994075, 60432289, 265006, 60433356, 60433438, 21906754, 87168474, 87168559, 265018, 264762, 264763, 264687, 21906765, 21906769, 27486262, 35695763, 18108374, 35696423, 264555, 18108385, 18108387, 18108388, 87168518, 264482
1513	87436228 (3025, 3026)	87436228 (3025, 3026) Novel Protein sim. GBank gi 1330394 (U58761) - C01F1.6 gene product [Caenorhabdilis elegans]			35696052, 264905, 264906, 264907, 264908, 264909, 264900, 264910, 264591, 264766, 264689, 264692, 264639, 264636

1514	95345392 (3027, 3028	1514 95345392 (3027, 3028) Novel Protein sim. GBank gil4559353]gblAAD23014.1IAC00658 - (AC006585) putative RIO1/2K632.3/MJ0444 family extragenic suppressor protein [Arabidopsis thaliana]	Contains protein domain (PF01163) - UNCLASSIFIED	UNCLASSIFIED	52644507, 52645156, 52646365, 52646842, 65274572, 22278994, 35696286, 56994075, 264259, 52645080, 29331822, 29331825, 269052, 2690562, 29331830, 52644045, 56182435, 265006, 60433356, 60433436, 52644296, 87168474, 87168559, 264448, 52644229, 21906765, 21906767, 21906768, 35695917, 265020, 52644150, 33657103, 52645129, 33657109, 33657109, 27486261, 27486262, 27486264, 27486265, 3669563, 18108376,
1515		79163536 (3029, 3030) Novel Protein sim. GBank gij3879501 jembjCAA87795 j - (Z47812) similar to ubiquitin carboxyl-terminal hydrolase; cDNA EST EMBL:D33366 comes from this gene; cDNA EST EMBL:D33965 comes from this gene; cDNA EST EMBL:D33822 comes from this gene; cDNA EST EMBL:D33822 comes from this gene; cDNA EST EMBL:D33822 comes from this gene; cDNA EST		ubiquitin	35996423, 35995855, 52644332, 18108385, 18108387, 87168518, 60432113, 265020, 264639
1516		88073539 (3031, 3032) Novel Protein sim. GBank gil498015 (L27479) - X123 [Homo sapiens] [Homo sapiens] 87793325 (3033, 3034) Novel Protein sim. GBank gil3415134 (AF082024) - Phyb1 [Pimpinella brachycarpa]		UNCLASSIFIED	265008, 56182323, 22279002 264091, 18108370, 264404
1518	87350697 (3035, 3036) 94328689 (3037, 3038)	87350697 (3035, 3036) Novel Protein sim. GBank gi728338 sp P39195 ALU8_HUMAN - !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! 94328689 (3037, 3038) Novel Protein sim. GBank gi15262681 emb CAB45771.1 - (AL080198) hypothetical protein [Homo sapiens]		tm7	66714117, 264508, 264509, 264905, 264510, 264910, 264910, 264591, 264595, 264288, 264766, 264769, 18108374, 264636, 264638, 264689, 264489, 60432049, 265009, 33657402, 264596, 21906754, 265019.
1520		87592855 (3039, 3040) Novel Protein sim. GBank gi 2662161 db BAA23712 - (AB007900) HH0452 cDNA clone for KIAA0440 has a 438-bp insertion at position 1711 of the sequence of KIAA0440. (Homo sapiens)		UNCLASSIFIED	264369, 21906765, 21906768, 21906769, 264691, 65274620, 33657182, 27486261, 18108374, 264557, 264639, 87168518, 22279002, 18108392, 60432049, 264259, 29331824, 265007, 60433356, 265010, 21906768, 264636
		86970696 (3041, 3042) Novel Protein sim. GBank gil5052351jgb AAD38516.1 AF13542 - (AF135421) GDP- mannose pyrophosphorylase B [Homo sapiens] 78960687 (3043, 3044)	Contains protein domain (PF00483) - synthase Nucleotidyl transferase		18108394, 264259, 66714117, 265011, 264603, 265019, 18108364, 35696423, 264557, 264558, 18108388
	91005151 (3045, 3046)	91005151 (3045, 3046) Novel Protein sim. GBank gij3776567 (AC005388) - Strong similarity to F21B7.33 gij2809264 from A. thaliana BAC gb[AC002560. EST gb[N65119 comes from this gene. [Arabidopsis thaliana]		UNCLASSIFIED	29331824, 265018, 265020, 265021 65274572, 21906768, 264693
1525	80203723 (3047, 3048) 87799867 (3049, 3050)	80203723 (3044, 3048) 87799867 (3049, 3050) Novel Protein sim. GBank gil4759040[ref NP_004283.1]pRIN1 - ras inhibitor		UNCLASSIFIED OUNCLASSIFIED	264112, 21906754, 263974 264683, 264687, 264689, 264690, 264692, 264693

15344 (3051 2052)	Mound Design of the Charle			
(2000), 0005)	95 1005 11 1	.	grycoprotein	35595285, 55182181, 60431735, 254595, 55812038, 264605, 264683, 21906765,
	S1/S2 PRECURSOR (GLUCAN 1,4-ALPHA-			55811957, 265020, 65274791, 264555,
	GLUCOSIDASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE)			264556, 264557, 264558, 264559, 83373044
(3053, 3054)	88262512 (3053, 3054) Novel Protein sim. GBank gi[2792496 (AF041107) - tuilp 2 [Rattus norvegicus]			56182575, 264259, 60432049, 29331822, 60432289, 264908, 66712502, 60433438, 87168559, 265017, 264288, 21906766, 21906769, 263977, 55811576, 568182323.
94130918 (3055, 3056)		5	UNCLASSIFIED	18108381 2227895, 22278997, 264259, 66712502,
-				264596, 265017, 265019, 264682, 264448, 264683, 264686, 21906765, 21906766, 21906768, 21906769, 265022, 264693, 83373044, 18108385
(3057, 3058)	94120793 (3057, 3058) Novel Protein sim. GBank gil4406663 gb AAD20053 - (AF131826) Unknown [Homo sapiens]	5	UNCLASSIFIED	264488, 263994, 56182575, 22278995, 35696286, 22278997, 264259, 29331822, 60432289, 29331827, 35696052, 264509,
				264906, 264907, 264908, 264909, 52644045, 56182435, 264511, 265009, 264910, 60433356, 60433438, 265017, 265018
				264760, 264448, 264764, 264369, 264288, 264768, 18108357, 264768, 52644229,
				21906765, 21906766, 21906767, 21906768, 265021, 265022, 52644150, 33657109,
				264629, 35695855, 60432113, 22279002, 264563, 264564, 264486, 264567
(3059, 3060)	95012765 (3059, 3060) Novel Protein sim. GBank gij2828710 (AF043642) - matrin cyclophilin [Rattus norvegicus]			264488, 264489, 35696286, 29331825, 35696052, 264508, 264905, 264907
				264909, 264510, 264511, 264512, 264910, 264502, 264505, 18108351, 264764, 264683
				264684, 264766, 264768, 18108357, 264769.
				35695917, 264628, 264629, 18108374, 35695855, 264630, 264631, 264634, 264555
				264636, 264637, 264404, 264563, 264566.
(3061 3062)	05.4.10354 (2061 2062) Novel Berton sim Charles (11005072 1100070)	THE STATE OF THE S		264486
(3001, 3004)	rover Frotein sin.; Spaint gij 19036/4 (US06/6) - Carboxyl terminal LIM domain protein (Homo sapiens)	Contains protein domain (PF00595) - kinase PDZ domain (Also known as DHR or	nase	561825/5, 35596286, 264097, 264259, 29331822, 29331825, 29331826, 29331827,
		GLGF).		35696052, 264509, 56182435, 264510,
				264511, 265007, 60433356, 55811386,
				264681, 264369, 264288, 264766, 264687,
				55811957, 35685817, 33657023, 35685763, 55810764 35696433 55811576 363981
				53816767, 53086723, 53811370, 205811, 60170394, 56182323, 83373044, 60432113.
				264566

264689	29331824, 29146499, 264907, 264112, 265008, 265011, 265017, 265018, 264762, 18108351, 263967, 20281149, 18108374, 263981, 264566	264905, 264907, 264766, 264637	62274572, 22278997, 264259, 60432049, 29331822, 60432289, 29331827, 29146499, 265006, 265008, 60170831, 60433438, 33109954, 87168559, 265018, 18108357, 21906788, 29148629, 265021, 265022, 18108377, 56182323, 60432113, 22279000, 22279002	264686, 18108357, 18108394, 21906767, 21906768, 29148629, 35696286, 265020, 265021, 52644150, 264693, 66714117, 29331825, 294693, 66714117, 20281149, 264909, 18108374, 35696423, 35695855, 265009, 264634, 264036, 264638, 18108385, 56526486, 265017, 265018, 264762, 18108351, 264448, 264369, 264766	65274572, 56182575, 60432049, 264259, 29331826, 265006, 265007, 60433356, 60433438, 264601, 18108351, 264448, 264369, 264288, 33657023, 65274620, 33657109, 60432113	18108394, 18108397, 264909, 265008, 265009, 265010, 18108351, 264638, 18108382, 18108385, 18108388
UNCLASSIFIED	struct	UNCLASSIFIED		Iranscriptfactor		kinase
				Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type		Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain
1532 85718224 (3053, 3064) Novel Protein sim. GBank gij3874716jembjCAA91265j - (Z66494) cDNA EST EMBL:D65271 comes from this gene; cDNA EST EMBL:D64845 comes from this gene; cDNA EST EMBL:D67438 comes from this gene; cDNA EST EMBL:D67438 comes from this gene; cDNA EST EMBL:D67638 comes from this gene; cDNA EST EMBL:D67638 comes from this gene; cDNA EST EMBL:D68087 comes from this gene; cDNA	94239830 (3065, 3066) Novel Protein sim. GBank gi 1490324 emb CAB01543 - (Z78141) unknown [Mus musculus]	95343941 (3067, 3068) Novel Protein sim. GBank gij81286 pir S22697 - extensin - Volvox carteri (fragment)	90936732 (3069, 3070)	1536 87602856 (3071, 3072) Novet Protein sim. GBank gi 106024 pir B32891 - finger protein 2, placental - human	95354556 (3073, 3074) Novel Protein sim. GBank gij3876332 embjCAB02096j - (Z79754) cDNA EST EMBL:T01054 comes from this gene: cDNA EST EMBL:D73600 comes from this gene; cDNA EST yk426f12.5 comes from this gene; cDNA EST yk342f10.5 comes from this gene; cDNA EST comes from this gene; cDNA EST comes from this gene; cDNA EST yk475c5.5	85724628 (3075, 3076) Novel Protein sim. GBank gi 403440 (MB1787) - [Gallus domesticus skeletal muscle mRNA, partial cds.], gene product [Gallus gallus]
1532	1533		1535			1538

22278994, 22278996, 35696286, 56994075, 22278998, 22278998, 22278999, 60432049, 264259, 29331822, 29331824, 60424269, 29331825, 60432289, 29331827, 29331826, 60432289, 29331827, 29331828, 35696052, 33656970, 56182435, 265009, 33657402, 60433356, 60433438, 55812038, 21906754, 87168559, 265017, 265018, 265019, 18108351, 264228, 21906764, 21906764, 21906764, 21906764, 21906764, 21906764, 35695917, 265020, 265021, 52644150, 33657023, 33657109, 27486261, 18108370, 18108376, 35696423, 55811576, 65274791, 264558, 56182323, 60170394, 83373044, 87168518, 60432113, 22279000, 264668	UNCLASSIFIED 264369, 264691, 263978	29331822, 35696052, 264259, 29331822, 35696052, 264508, 264509, 264905, 264907, 264908, 264501, 264509, 264907, 264907, 264908, 264501, 264509, 264701, 264761, 264761, 264764, 264288, 264687, 21908769, 55811957, 35695917, 265020, 264691, 33657023, 264692, 33657109, 264638, 18108374, 264632, 264636, 264636, 264868	cathepsin 18108392, 18108394, 65274572, 29331822, 264508, 265007, 265008, 265009, 265011, 264682, 18108354, 18108355, 52644150, 18108368, 264636, 18108381, 18108382		id_recpt 35696286, 56994075, 22278999, 35696052, 60433356, 60433438, 265011, 264683, 33657109, 35696423, 264631, 87168518, 22279000		UNCLASSIFIED 264905, 264686 UNCLASSIFIED 264259, 29331822, 66714117, 265007, 55811386, 285010, 264600, 265017, 265019, 264288, 264788, 265020, 265022, 55811576,
Ē	5	1a	90		Contains protein domain (PF00439) - nuci_recpt Bromodomain	5	5.5
95337628 (3077, 3078) Novel Protein sim. GBank gij3218411 jembjCAA19575.1] - (AL023659) SPBC19C7.07c, putative tRNA splicing endonuclease ga mma subunit, len:284aa, similar eg. to YAR008W, YAHB_YEAST, P39707, YAR008W, tRNA splicing endonuclease gamma subunit, (275aa), fasta scores, opt:269, E():6.4e-2	95352858 (3079, 3080) Novel Protein sim. GBank gi 5052634 gb AAD38647.1 AF14567 - (AF145672) BCDNA.GH12174 Drosophila melanogaster]	95317948 (3081, 3082) Novel Protein sim. GBank gi 5052349 gb AAD38515.1 AF13501 - (AF135016) protein phosphatase 2A 48 kDa regulatory subunit [Homo sapiens]	90937549 (3083, 3084) Novel Protein sim. GBank gi[5305702]gb AAD41779.1 AF12686 - (AF126867) calpain- like protease [Mus musculus]	84348768 (3085, 3086) Novel Protein sim. GBank gi 728832 sp P39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!	1544 87757295 (3087, 3088) Novel Protein sim. GBank gil3493162 (AF084259) - bromodomain-containing protein BP75 [Mus musculus]	85757973 (3089, 3090) Novel Protein sim. GBank gil1086591 (U41007) - similar to S. cervisiae nuclear protein SNF2 (SP:P22082) in a a region of giy-arg repeats (Caenorhabditis elegans)	86999594 (3093, 3094) Novel Protein sim. GBank gil2661132 (AF035683) - p21 [Mus musculus]
9 95337628 (3077, 3078)	95352858 (3079, 3080)	95317948 (3081, 3082)	90937549 (3083, 3084)	1543 84348768 (3085, 3086)			1547 86995594 (3093, 3094)

1548	84233065 (3095, 3096)	1548 94233065 (3095, 3096) Novel Protein sim. GBank gil3043692 dbj BAA25510 -		UNCLASSIFIED	29331824, 60431528, 264639, 56182323
5249		95330048 (3097, 3098) Novel Protein sim. GBank gil569519 dbj BAA83043.1 - (AB029014) KIAA1091 protein [Homo sapiens]		136. 37.	60424179, 22278995, 35696286, 22278998, 22278998, 22278999, 264092, 264094, 29331822, 56182181, 29331824, 35686052, 264905, 264908, 264909, 265006, 264511, 265008, 60431735, 60433356, 21906754, 5581136, 8168559, 265017, 265018, 265019, 55811150, 264682, 264288, 264369, 21906769, 51906769, 21906769, 21906769, 21906769, 21906769, 21906769, 215626486, 60432113, 22279002, 264563, 265668
		95201907 (3099, 3100) Novel Protein sim. GBank gi 544463 sp P35350 GUSB_BOVIN - POSSIBLE GUSTATORY RECEPTOR TYPE B (PPR1 PROTEIN)	Contains protein domain (PF00001) - tm7 7 transmembrane receptor (rhodopsin family)		65274572, 60432289, 265008, 264910, 265011, 265017, 265019, 264768, 56182323
		88077111 (3101, 3102) Novel Protein sim. GBank gil4756566[ref NP_004798.1 pHS6S - heparan-sulfate 6- sulfotransferase		UNCLASSIFIED	22278999, 29331822, 264508, 264509, 264906, 264907, 264909, 265007, 264512, 264910, 21906754, 265018, 265019, 264681, 264764, 264766, 264688, 26469, 21906769, 264692, 35695763, 264635, 264555, 264556, 264657, 26457, 2645
	87617114 (3103, 3104)			UNCLASSIFIED	264259, 29331828, 66712502, 264764, 264288, 264686, 33657109, 264556
	94725512 (3105, 3106) ,		Contains protein domain (PF00304) - dehydrogenase Gamma-thionins family		56182575, 35696286, 29146499, 264509, 264907, 265908, 264908, 264909, 56182435, 265006, 265006, 265009, 264910, 264757, 264758, 265017, 55811150, 18108351, 2645744, 56181562, 35695917, 264693, 33657109, 18108374, 35696423, 65274791, 35695855, 264635, 264555, 56182323, 18108382,
1554	94233069 (3107, 3108)	94233069 (3107, 3108) Novel Protein sim. GBank gi]3043692[dbj]BAA25510] - (AB011156) KIAA0584 protein [Homo sapiens]	Contains protein domain (PF00446) - transferase Gonadotropin-releasing hormones		35696286 . 22278997 . 264259, 29331822. 29331824 . 255007 . 265009 . 60432229 . 33657402 . 55812038 . 265011 . 265019 . 264681 . 264369 . 264686 . 264767 . 264768 . 21906765 . 21906769 . 3569517 . 264693 . 18108370 . 80431528 . 55811576 . 264631 . 60170394 . 56182323 . 83373044 . 18108385 . 22279000 . 22279002

264259, 29331826, 35696052, 264508, 264905, 264906, 264907, 264908, 52644045, 264909, 264900, 264900, 264900, 264900, 264900, 264900, 264900, 264900, 264900, 265010, 8716859, 264759, 33657084, 264763, 264764, 264268, 264766, 264768, 21906765, 3569585, 264631, 33657023, 35695855, 264635, 264586, 264555, 264536, 264586, 264555, 264636, 264638	83373044, 264758, 265022, 264600, 35696052, 264630, 35696423, 265018, 264632, 264682, 29331822, 265020, 265011	60432289, 264509, 264906, 264907, 264908, 264909, 264909, 264910, 264758, 55811388, 264761, 264762, 264769, 264690, 263978, 264634, 264635, 264539, 264564, 264486	22278994, 22278996, 22278997, 22278998, 22278999, 60432049, 264259, 29331824, 29331825, 29331827, 264908, 264909, 6043356, 21906754, 265017, 265018, 26448, 21906767, 265017, 265018, 26448, 21906767, 265021, 26502, 33657109, 18108370, 55811576, 83373044, 87168518, 22279000, 22279002	264259, 29331622, 60432289, 35696052, 264107, 264110, 21906754, 33109954, 87168559, 264760, 264763, 21906764, 21906765, 21906769, 265021, 264690, 35695855	264908, 264603, 264638	264510, 264594 263967	264910, 264764, 264766	18108394, 35696286, 264259, 29331822, 60432289, 35696052, 29331828, 264508, 66712502, 264908, 56182435, 265007, 264910, 60170831, 21906754, 265011, 265017, 265018, 265019, 18108351, 26448, 264288, 264686, 21906765, 21906768, 265021, 60170615, 264692, 35696423, 35695855, 264557, 56182323, 60432113, 222730002, 264482
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		glycoprotein	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	dehydrogenase
Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat		Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)		Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat				Contains protein domain (PF00441) - dehydrogenase Acyl-CoA dehydrogenase
1555 87332970 (3109, 3110) Novel Protein sim. GBank gij2257495 dbj BAA21392 - (AB004534) pi015 [Schizosaccharomycas pombe]	ic	87640609 (3113, 3114) Novel Protein sim. GBank gil3329611 (AF078783) - contains similarity to C3HC4-type zinc fingers (Pfam; zf-C3HC4-tym, score; 34.08); most similar to drosophila goliath protein (SW: Q06003) [Caenorhabditis elegans]	94840376 (3115, 3116) Novel Protein sim. GBank gi[5360105]gbJAAD42871.1[AF15510 - (AF155105) putative zinc finger protein NY-REN-34 antigen [Homo sapiens]	88224865 (3117, 3118) Novel Protein sim. GBank gij112908 spjP02750 A2GL_HUMAN - LEUCINE-RICH ALPHA-2-GLYCOPROTEIN (LRG)	84580675 (3119, 3120) Novel Protein sim. GBank gij3880146 emb CAA92704 - (Z68319) Similarity to Human hnRNP F protein (PIR Acc. No. S43484); cDNA EST EMBL:D34218 comes from this gene; cDNA EST EMBL:D37248 comes from this cDNA EST EMBL:D71817 comes from this gene; cDNA EST EMBL:D71817 comes from this gene; cDNA	2)		87766371 (3127, 3128) Novel Protein sim. GBank gi[1168287]sp[P45953]ACDV_RAT - ACYL-COA DEHYDROGENASE, VERY-LONG-CHAIN SPECIFIC PRECURSOR (VLCAD)
6 87332970 (3109, 3110	91229268 (3111, 3112)						3 85508694 (3125, 3126)	
155	1556	1557	1558	1559	1560	1561	1563	1564

4/51	94216142 (3147, 3148)	94216142 (3147, 3148) Novel Protein sim. GBank gil4758334 ref NP_004256.1 pFADS - delta-6 fatty acid desaturase	Contains protein domain (PF00173) - cytochrome Heme-binding domain in cytochrome b5 and oxidoreductases	сутостате	18108394, 264887, 18108397, 18108398, 22278996, 22278999, 22278999, 264259, 29331825, 29346498, 29146499, 2654107, 264907, 264909, 52644045, 265511, 265019, 18108351, 264682, 264763, 264764, 18108354, 264388, 264688, 264686, 21906766, 21906766, 21906767, 21906768, 21906769, 2
1575		95340019 (3149, 3150) Novel Protein sim. GBank gil3881810 emb CAA94856 - (Z70783) similar to EF-hand calcium binding protein; cDNA EST EMBL:C08700 comes from this gene [Caenorhabditis elegans]	Contains protein domain (PF00036) - phosphatase EF hand	phosphalase	56994075, 264259, 29331822, 29331824, 29331825, 60432289, 29331828, 264909, 265006, 265008, 265010, 87168559, 55811150, 264448, 18108354, 264369, 265021, 60170615, 264691, 33657023, 33657109, 60431528, 65274791, 35695855, 18108385, 60432113, 22279002, 264482
1576		95314019 (3151, 3152) Novel Protein sim. GBank gi[2773195 (AF039711) - contains similarity to Physcomitrella patens glyceraldehyde 3-phosphate dehydrogenase (GB:X72381) [Caenorhabditis elegans]	-	UNCLASSIFIED	264569, 264092, 264094, 264095, 264259, 264508, 264508, 264509, 264909, 264511, 265006, 264910, 21906754, 265010, 265011, 87168559, 264761, 264288, 264766, 264761, 264288, 264693, 264691, 264693, 35695855, 264632, 264634, 264636, 264638, 264688,
1577	87613800 (3153, 3154) 87123138 (3155, 3156)	87613800 (3153, 3154) Novel Protein sim. GBank gi[2499130]sp[P70315]wASP_MOUSE - WISKOTT- ALDRICH SYNDROME PROTEIN HOMOLOG (WASP) 87123138 (3155, 3156)		UNCLASSIFIED	264511, 265011, 264681, 264369, 264686, 264689, 264629, 264555, 264558, 264559 264259, 29331826, 265017, 264689, 264693,
1579		88085141 (3157, 3158) Novel Protein sim. GBank gij2978255[dbjjBAA25190] - (AB007407) myeloid zinc finger protein-2 [Mus musculus]	Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	UNCLASSIFIED	60432113 35696286, 264908, 264909, 60433438, 55811386, 264369, 264685, 33657023, 264555, 264556, 264557, 87168518
1580		87255702 (3159, 3160) Novel Protein sim. GBank gil4324682 gb AAD16986 - Contains protein domain (PFG (AF109674) late gestation lung protein 1 [Rattus norvegicus] SCP-like extracellular protein 95087431 (3161, 3162) Novel Protein sim. GBank gi 2088838 (AF003386) - F59E12.4 gene product [Caenorhabditis elegans]	Contains protein domain (PF00188) - glycoprotein SCP-like extracellular protein	glycoprotein	22278999, 35696052, 29331830, 52644045, 55812038, 87168474, 265018, 264448, 265022, 264638, 56526486, 22279000 22278995, 29331822, 29331824, 29331826, 56182435, 264595, 55812038, 87168559, 265017, 264288, 21906764, 55811957, 35695917, 284692, 55811576, 264637,
					56182323, 264559, 83373044, 60432113

264259, 60432289, 29331827, 264509, 264905, 264900, 264900, 264910, 264762, 264288, 264768, 264769, 264632, 264555, 264639, 56526486, 22279000	60170831, 33657402, 264682, 21906766. 35695855, 264563	60424179, 52646842, 65274572, 56182575, 22278998, 22278996, 252278998, 22278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 25278999, 254526, 29331827, 25584045, 56182435, 264510, 224512, 265008, 6043336, 33657402, 60433438, 25812038, 21906754, 55811386, 52644296, 87168474, 87168559, 265018, 265019, 266448, 264369, 265018, 265020, 265027, 265022, 25264150, 33657023, 33557109, 18108374, 55810754, 56182323, 83373044, 18108387, 87168518, 60432113, 22279002	35696286, 22278998, 264259, 29331822, 29331824, 29331825, 264905, 265006, 265007, 265008, 60433356, 33109954, 87168474, 265011, 265017, 264604, 264369, 264288, 264685, 264769, 18108359, 21906765, 18108364, 18108388, 264482, 264564	265017, 265018, 264689, 33657023, 263978, 264636, 264563	264907, 264908, 264511, 264910, 264591, 264594, 264629, 264631, 264563, 264483, 264567	264259, 29331828, 264905, 265006, 264758, 21906754, 264761, 264762, 21906765, 21906769, 60170615, 52644150, 33657109, 35695855, 56182323, 18108385	65274572, 264490, 29331822, 66714117, 29331827, 29331828, 56182435, 265008, 60170831, 264595, 264758, 264596, 265011, 264686, 21906766, 21906768, 55811957, 27486265, 264639, 18108385, 56526486, 60432113
phosphalase	ÜNCLASSIFIED	dehydrogenase	phosphatase		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase	Contains protein domain (PF00023) - phosphatase Ank repeat			Contains protein domain (PF00468) - Ribosomal protein L34	
1582 95358052 (3163, 3164) Novel Protein sim. GBank gi 5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Lelshmania major]	87622715 (3165, 3166) Novel Protein sim. GBank gil5578958 emb CAB51351.1 - (AL050306) dJ47587.2 (novel protein) [Homo sapiens]	95337722 (3167, 3168) Novel Protein sim. GBank gil5531815 gblAAD4482.1 - (AF078850) steroid dehydrogenase nomolog [Homo sapiens]	87626117 (3169, 3170) Novel Protein sim. GBank gil4240132 dbj BAA74846.1 - (AB020630) KIAA0823 protein [Homo sapiens]	88067081 (3171, 3172) Novel Protein sim. GBank gij3786494 (AF098993) - No Idefinition line found [Caenorhabditis elegans]	87617126 (3173, 3174) Novel Protein sim. GBank gi[3253159 (AF005355) - translation initiation factor eIF2C [Oryctolagus cuniculus]	87802536 (3175, 3176) Novel Protein sim. GBank gi/1077573/pir/I S52680 - probable Contains protein domain (PF00468) - UNCLASSIFIED ribosomal protein L34 mitochondrial - yeast (Saccharomyces cerevisiae)	90980653 (3177, 3178) Novel Protein sim. GBank gi[2137756 pir 48746 - semaphorin C - mouse (fragment)
95358052 (3163, 3164)	87622715 (3165, 3166)	95337722 (3167, 3168)					
1582	1583	1584	1585	1586	1587	1588	1589

1590				UNCLASSIFIED	264489, 22278996, 264259, 29331824, 29331825, 29331825, 29331826, 29331827, 265006, 60433356, 21906754, 265017, 265018, 265019, 264448, 264765, 264288, 5264429, 21906765, 21906765, 21906769, 265021, 264692, 2746256, 35695763, 265021, 264692, 2746256, 35695763, 2652486, 61437113, 22779010, 22729010
1591					264564
1592		97882533 (3183, 3184) Novel Protein sim. GBank gil4557749 ref NP_000237.1 pMHC2 - MHC class II iransactivator		MHC	264259, 264905, 29331830, 264595, 265017, 264448, 264288, 264690, 264629, 87168518
1593				UNCLASSIFIED	65274572, 60432049, 264509, 60433356, 21906754, 21906767, 21906768, 18108370, 3469423, 22270000, 284862, 284867
1594		87773752 (3187, 3188) Novel Protein sim. GBank gij3877072jembjCAA87060j - (726937) similarity with ribectural protein 1.31		UNCLASSIFIED	264488, 29331827, 264905, 264906, 264907.
		[Caenorhabditis elegans]			264908, 264909, 264910, 264592, 264593, 264757, 264602, 264604, 264760, 264681, 264288, 264768, 264769, 264881,
					20450, 204100, 204100, 23140023, 35095917, 264692, 264628, 264629, 264630, 264632, 264634, 264635, 264638, 264639, 204563, 264634, 264638, 264639,
1595		2703 (AF065389) -	Contains protein domain (PF00335) - UNCLASSIFIED	UNCLASSIFIED	29331826, 264908, 55811957
		כייסיקטיין יידי די דייסיויס אמקופונאן	4 transmembrane segments integral membrane proteins		
8	_			SSIFIED	29146498, 264758, 263967
1597		65971857 (3193, 3194) Novel Protein sim. GBank 1997 (3193, 3194) 1995257114 1997 (4F094480) 1997 (4F094480) 1998 (4F094480) 1998 (4F094480) 1998 (4F094480) 1998 (4F09460804888 Homo saniens) 1998 (4F094888 Homo saniens) 1998 (4F09488 Homo saniens) 1998 (4F094888 Homo saniens) 1998 (4F09488 Homo saniens) 1998 (4F0	Contains protein domain (PF00067) - cyto450 Cytochrome P450		264092, 29331824, 264508, 264682, 264369, 264686, 264630, 264563
1598					264259 264634
1599		- 5	Contains protein domain (PF01581) - UNCLASSIFIED		52645080, 29331824, 29331826, 264511,
		girəco əz irelivr 7 (olivopontocerebellar atrop	Ipocar - spinocerebelar ataxia rmkramide related peptide family hy with retinal degeneration)		265009, 265011, 264605, 264448, 264764, 265020, 264692, 264693, 18108370, 264635, 18108385
8	$\overline{}$				29331826, 264603, 264691, 264563
160	_	15023246 (3201, 3202)		ASSIFIED	264635
7091		1 AF12853 - (AF128535) in PACSIN2 [Mus musculus]	Contains protein domain (PF00018) - struct SH3 domain		29146499, 264112, 264762, 18108351, 29148627, 263974
1603		Novel Protein sim. GBank gi 283920 pir S27939 - Iensin - chicken		collagen	264490, 29331824, 264907, 264909, 264511, 265008, 264592, 265010, 265011, 264762, 264764, 264369, 264288, 264687, 264769, 264693, 264628, 264636, 264636, 264556, 264556, 264558, 264558, 264559, 264558, 264559, 264559,
1604	80221813 (3207, 3208)	80221813 (3207, 3208) Novel Protein sim. GBank gil4768831[gb[AAD29633.1]AF11682 - (AF116827)		ATPase_associated	18108385 263977
		unknown [Homo sapiens]			

1605	1605 91221129 (3209, 3210)			struct	264905, 264509, 264906, 264907, 264908,
					264909, 264604, 264766, 264768, 264692,
					264693, 33657109, 264629, 35695855. 264635, 264636, 264637
1606	94312703 (3211, 3212)	94312703 (3211, 3212) Novel Protein sim. GBank gil4505313jref NP_003794.1pMYOM - UNKNOWN	Contains protein domain (PF00047) - struct Immunoglobulin domain	struct	22278996, 22278999, 264259, 33657402, 265017, 18108351, 264448, 21906767, 21906769, 52644150, 264691, 87168518
1607	10871805 (3213, 3214)	10871805 (3213, 3214) Novel Protein sim. GBank gi[5174473]ref[NP_005888.1 piPP] - intracisternal A particle- promoted polypeptide		transcriptfactor	264689
1608		80428900 (3215, 3216) Novel Protein sim. GBank gi[2224629]dbj BAA20802 - (AB002342) KIAA0344 [Homo sapiens]		UNCLASSIFIED	264094, 264906, 264907, 264909, 264910, 264591, 264603, 264768, 264693, 264634, 264635, 264637, 264639
1609		94311572 (3217, 3218) Novel Protein sim. GBank gil4884073 emb CAB43213.1 - (AL.049934) hypothetical protein [Homo sapiens]	·		52644507, 52645156, 52646365, 52646842, 56182575, 22278994, 56994075, 35696286, 22278997, 22278998, 22278999, 264259,
					52645080, 29147620, 29331826, 35696052, 33656970, 264508, 264509, 264907, 52644045, 56182435, 264510, 264511,
					264512, 33657402, 21906754, 52646317, 33109954, 52644296, 87168474, 265017, 265018, 265019, 18108351, 264448, 264288,
					264769, 52644229, 21906765, 21906766, 21906767, 21906767, 21906768, 21906769, 55811957.
					35695917, 265020, 265021, 265022, 52644150, 33657023, 33657109, 52645129,
					27486261, 27486262, 35695763, 264628, 18108370, 18108376, 35696423, 264638, 52644332, 18108387, 87168518, 22279000,
1610	85468200 (3219, 3220)	85468200 (3219, 3220) Novel Protein sim. GBank gi[283920 pir 527939 - tensin -		UNCLASSIFIED	264563, 264486 264593, 264757, 55812038, 265018, 265020, 264691, 264692, 264633, 264634
		chicken			264635, 264555, 22279000, 264564
191	94122843 (3221, 3222)	94122843 (3221, 3222) Novel Protein sim. GBank gi 107284 pir A35415 - Contains protein peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (fragment)	Contains protein domain (PF00008) - peroxidase n EGF-like domain	peroxidase	35696286, 21906765, 264691, 35696423
1612		85746031 (3223, 3224) Novel Protein sim. GBank gil3874846 emb CAA94337 - (270307) Similarity to B.subtilis tetracycline resistance protein (SW/TCR2_BACSU); cDNA EST EMBL: C09951 comes from this name: cDNA EST EMBL: C08265 comes		UNCLASSIFIED	264486, 264509, 18108370, 18108387, 264486
					CJETOG
1613	82247354 (3225, 3226)	_		UNCLASSIFIED	264759

<u> </u>	91220634 (3227, 3228,	1014 91zz6654 (3zz7, 3zz6) Novel Protein sim. Gbank di4680673 qb AAD27726.1 AF13295 - (AF132951) CGI-17	Contains protein domain (PF01605) - UNCLASSIFIED PRF1-like proteins		22278995, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 264908
					264512, 265009, 265011, 265017, 265018,
					265019, 18108351, 264683, 264288, 264766,
					21906767, 21906768, 21906769, 35695917.
		-			265021, 265022, 35696423, 35695855,
١					60170394, 56182323, 83373044, 264566
1615		86121909 (3229, 3230) Novel Protein sim. GBank gij5689485 dbj BAA83026.1 -	Contains protein domain (PF00023) - homeobox		22278996, 35696286, 22278997, 29331822,
		(AB028997) KIAA1074 protein [Homo sapiens]	Ank repeat		35696052, 29331828, 264508, 264908,
					264909, 56182435, 264511, 265017, 265019,
					264766, 264767, 264768, 265020, 264691,
		-			264628, 264632, 264635, 264555, 264556,
	_				56182323, 264558, 22279002
1616				UNCLASSIFIED	264488, 52644507, 52645156, 52646365,
		(Z78418) cDNA EST EMBL:D71020 comes from this gene;			52646842, 22278994, 22278995, 35696286,
		cDNA EST EMBL:D73593 comes from this gene; cDNA			22278996, 22278997, 22278999, 52645080,
		EST EMBL: C07649 comes from this gene; cDNA EST			29331822, 29331824, 29331825, 29331827,
		EMBL:C09081 comes from this gene; cDNA EST yk399f2.3			29331828, 35696052, 33656970, 264905.
		comes from this gene; cDNA			264909, 264594, 52646317, 21906754,
					33657084, 52644296, 87168474, 87168559,
					265017, 265018, 265019, 264681, 264448,
					264684, 52644229, 21906764, 264689,
					21906765, 21906766, 21906769, 35695917,
					265020, 265021, 52644150, 33657023,
					52645129, 33657109, 33657182, 27486261.
					27486262, 33657349, 27486265, 35695763,
					18108376, 35696423, 35695855, 264557,
					52644332, 264558, 18108385, 87168518
1617	_	88090742 (3233, 3234) Novel Protein sim. GBank	Contains protein domain (PF01529) - peptidase		35696052, 264905, 264509, 264907, 264908
	_	gil466053 sp P34679 YO41_CAEEL - HYPOTHETICAL 68.7 DHHC zinc finger domain	DHHC zinc finger domain		264510, 264511, 264764, 264766, 264768,
_		KD PROTEIN 2K757.1 IN CHROMOSOME III)		264689, 264693, 18108374, 264635, 264636,
	_				264638
1618		86272860 (3235, 3236) Novel Protein sim. GBank gil4240231 dbjjBAA74894.11 -		struct	35696286, 22278999, 264092, 29331824,
		(AB020678) KIAA0871 protein [Homo sapiens]			29331825, 35696052, 33657084, 21906765,
					27486264
1619		95354580 (3237, 3238) Novel Protein sim. GBank	Contains protein domain (PF00010) - Iranscriptfactor	transcriptfactor	52646842, 65274572, 22278999, 264259,
		gij5031763 ref NP_005515.1 pHRY - hairy (Drosophila)-	Helix-loop-helix DNA-binding domain		29331822, 29331824, 29331825, 29331826,
		homolog		-	29331827, 29331828, 35696052, 56182435,
					265007, 265008, 264910, 60170831,
					60432229, 60433356, 60433438, 265019,
					264448, 264288, 264686, 21906768, 265021,
					60170615, 33657023, 65274620, 33657109,
					18108374, 18108376, 35696423, 35695855,
					56182323, 56526486
1620	87344655 (3239, 3240)	1620 87344655 (3239, 3240) Novel Protein sim. GBank		UNCLASSIFIED	264684
		PROTEIN			

1634	87076700 /2244 2245				
1622		Mount Design of Contract of the Contract of th		UNCLASSIFIED	264910
		Calmodulin-binding protein [Gallus gallus] SPRY domain (PF00622) - UNCLASSIFIED	r Contains protein domain (PF00622) - SPRY domain	UNGLASSIFIED	18108392, 65274572, 18108398, 22278996, 22278997, 22278999, 29146498, 29146499, 264905, 264908, 264909, 264908, 29146499, 264905, 264908, 264909, 264908, 264909, 264909, 264909, 264909, 264909, 265017, 264389, 21906758, 58811957, 265021, 60170615, 264635, 264557, 60170039, 83373044, 18108385, 22279000, 267270000, 264668
1623	87779106 (3245, 3246)	87779106 (3245, 3246) Novel Protein sim. GBank gil731086jsp P40389 UV22_SCHPO - UV-INDUCED PROTEIN UVI22		ribosomalprot	18108398, 264250, 264909, 56182435, 18108398, 264248, 21906768, 35695917, 26469, 2166958, 35695917, 264691, 2166958, 35695917, 264691, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 2646918, 2166958, 264998, 2166958, 264998, 2166958, 264998, 2166958, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 2166658, 216665
1624		87338178 (3247, 3248) Novel Protein sim. GBank gij3875666jemb CAB05478] - (Z83104) cDNA EST EMBL:100015 comes from this gene; cDNA EST EMBL:D33665 comes from this gene; cDNA EST EMBL:D36540 comes from this gene; cDNA EST yk240f8.3 comes from this gene; cDNA EST yk387c8.3 comes from this gene; cDNA EST yk387c8.3		UNCLASSIFIED	264758
1628		94734369 (3251, 3252) Novel Protein sim. GBank gil489622 dbjjBAA76833.11 - (AB023206) KIAA0989 protein [Homo saplens] 94734369 (3251, 3252) Novel Protein sim. GBank gil5679070[gb]AAD46844.1 JAF16090 - (AF160904) BCDNA. HL05936 [Drosophita melanogaster]		kinase	264489, 22278994, 22278995, 22278996, 35696286, 22278997, 22278998, 22278999, 224092, 264092, 264289, 29331824, 29331825, 29331824, 29331825, 29331827, 29331824, 29331825, 29331827, 29331824, 264106, 264508, 3857084, 265017, 265018, 18108351, 264681, 264369, 65274620, 35695917, 265021, 264691, 65274620, 35695917, 265021, 264691, 65274620, 25278994, 26279000, 22279002, 22278994, 26245156, 52646365, 52646842, 22278999, 264289, 26278999, 264289, 26278999, 264289, 26278999, 264331822, 29331824, 6671417, 29331826, 60432289, 26331824, 6643229, 264409, 2643948, 264389, 264489, 264389, 264489, 264389, 264489, 264389, 264486, 265014, 265022, 264406, 265018, 264449, 264502, 27486261, 27486262, 27486261, 27486262, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486262, 27486261, 27486261, 27486261, 27486262, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486261, 27486262, 27486261, 27486262, 27486261, 274
1627	83368773 (3253, 3254)			UNCLASSIFIED	264288
1628	85708459 (3255, 3256)	85708459 (3255, 3256) Novel Protein sim. GBank gij3668087 (AC004667) - Invoorhetical protein Azabidonsis thatianal	8		264288, 264686, 264767, 22279002
		Thomsered Protein (Nationalists trialiana)			

1915692 emb CAA69995 - In [Homo sapiens] 1915692 emb CAA69995 - Ins gallus] F15075 - (AF150755) F15075 - (AF150755) F15075 - (AF150765) Contains protein domain (PF00435) - Spectrin repeat factor [Mus musculus] Contains protein domain (PF0069) - DAPK - death-associated factor [Aco04680) - Ithaliana] 420389 emb CAB46680.1 - Ican [Leishmania major]	emb[CAA69995 - stylens] emb[CAA69995 - ctor [Mus musculus] (AF150755) Spectrin repeat AC004680) - cmb[CAB46680.1 - cmb[CAB46680.1 - cmb]CAB46680.1 - cmb]CAB46680.1	264555	UNCLASSIFIED 29331822, 29331827, 265010, 264693, 264534, 22278002	UNCLASSIFIED 55811957, 264259, 33657023, 264693,	29331822, 29331824, 29331827, 29331828, 264906, 264908, 55811576, 264910, 264634,	264636, 264637, 56182323, 264559, 264758,	UNCLASSIFIED (264687, 264769, 264691, 264692, 29146499		264988, 264509, 264907, 264909, 264909,	265007 264637 22279002		UNCLASSIFIED 65274572, 22278998, 35696052, 52644045,	264511, 265008, 265009, 265010, 265011,	265018, 265019, 264448, 264369, 21906765,	INC. ACCIETED 2007800 PEACON 201700 ACCIETED	Solrico	е 264558	21906765, 21906767, 22278996, 35696286	22278999, 264259, 264692, 264693,	29331824, 33657109, 264508, 264906,	18108370, 264629, 265007, 33657402,	21906754, 264602, 264604, 264764, 264683. 264566, 264288	UNCLASSIFIED 284488, 18108394, 65274572, 56182575.	35696286, 29331824, 29331826, 29331827, 35696052, 264908, 56182435, 264112	265006, 265008, 264757, 264758, 55811386	264603, 264760, 18108351, 264764, 264288,	264766, 264768, 21906767, 55811957,	[264691, 33657023, 65274620, 18108370]	
1240175 dbj BAA74866.1 - 1 [Homo sapiens] 1915892 emb CAA69995 - 1 [Homo sapiens] 1915892 emb CAA69995 - 1 [Homo sapiens] 1915851 dbj BAA22896 - 1 [Homo sapiens] 1915889 emb CAB46680 - 1 [Homo sapiens] 19158889 emb CAB46680 - 1 [Homo sapiens] 1915889 emb CAB46680 - 1 [Homo sapiens] 1915889 emb CAB4680 - 1 [Homo sapiens] 1915	(AB020650) KIAA0843 protein [Homo sapiens] (Y08779027 (3259, 3264) Novel Protein sim. GBank gil2558501[dbj]BAA22896] - (Y08740) tom-1A protein [Gallus gallus] (Y08740) tom-1A protein [Gallus gallus] (Y08740) tom-1A protein [Gallus gallus] (Y08740) tom-1A protein sim. GBank gil2558501[dbj]BAA22896] - (D83850) hepatoma-derived growth factor [Mus musculus] (Y08773683 (3265, 3266)) Novel Protein sim. GBank gil2558501[dbj]BAA22896] - (AF150755) Microtubule-actin crosslinking factor [Mus musculus] (Y08740) Novel Protein sim. GBank gil3420051 (AC004680) - unknown protein [Arabidopsis thaliana] (AJ243460) protein sim. GBank gil5420389[emb]CAB46680.1 - (AJ243460) protein sim. GBank gil5420389[emb]CAB4680.1 - (AJ243460) protein sim. GBank gil54203889[emb]CAB4680.1 - (AJ243460) protein si	struc	ONO	ONO			ONO			Contains protein domain (PF00435) - Istruc	Spectrin repeat	ONO				ONO	Contains protein domain (PF00069) - kinas Eukaryotic protein kinase domain						ONO						
	87779027 (3257, 3258 87779027 (3259, 3260 87773683 (3263, 3264 87871692 (3263, 3264 87871692 (3263, 3266 85992817 (3267, 3268) 80070435 (3271, 3272) 80070435 (3275, 3276) 87101854 (3275, 3276)	NOVEL FIOLEN SIM. GBank glt4240175 db BAA/4856.1 - (AB020550) KIAA0843 protein [Homo sapiens]		-			Novel Protein sim. GBank gi[2558501 dbj BAA22896 -	(D63850) hepatoma-derived growth factor [Mus musculus]			gil4887229 gb AAD32244.1 AF15075 - (AF150755) mlcrotubule-actin crossiinking factor [Mus musculus]						7.1pDAPK - death-associated	Novel Protein sim. GBank gij3420051 (AC004680) -	unknown protein [Arabidopsis thaliana]				Novel Protein sim. GBank gi[5420389 emb CAB46680.1 -	recovery proceedingsyrail [Leisimaina major]					

56182575, 56994075, 35696286, 60432049, 60432289, 29331827, 35696052, 52644045, 56182435, 264510, 265006, 265007, 265008, 264910, 265009, 3365702, 55812038, 265010, 265011, 265017, 265018, 264282, 21906765, 21906766, 21906768, 35695917, 265021, 60170015, 52644150, 33657023, 33657109, 33657349, 18108374, 35696423, 65274791, 35695855, 264632, 264555, 56182323, 22279000	29146499, 265006, 265007, 265008, 265009, 55812038, 265010, 265011, 264555, 264556, 264558, 18108383	22278999, 29147620, 29331826, 29331828, 33656970, 55812038, 265010, 265018, 265019, 18108351, 264689, 265020, 265022, 264630, 33657023, 65274620, 35695763, 52644332, 18108381, 60170394, 56182323, 18108388, 87168518, 22279002, 264564	29331825, 29331827, 29331828, 21906754, 265019, 264288, 264693, 33657349, 18108370, 18108376, 264555, 83373044, 22279002, 264482	264905, 264907, 264908, 264910, 265009, 264757, 264758, 264761, 264762, 264763, 264766, 264628, 264629, 264630, 264507, 264565, 264566, 264565, 264566, 264566, 264565, 264566, 264565, 264567, 264567, 264567, 264567, 264568	264685, 264693	22278994, 56994075, 35696286, 264259, 29331824, 29331825, 29331826, 60432289, 264508, 60433356, 60433438, 87168559, 265018, 264687, 35695917, 264692, 33657182, 27486261, 27486265, 33657349, 60432113, 264564	29331822, 264906, 264908, 264369, 21906768, 60170615, 264639, 22279000	265009, 264686, 55811957, 35695917, 55810764, 264556, 56182323, 264558, 18108385
UNCLASSIFIED	UNCLASSIFIED	struct		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF01843) - struct DIL domain						Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type
1640 94143185 (3279, 3280) Novel Protein sim. GBank gi]2842469 emb CAA16847.1 - (AL021747) hypothetical protein {Schizosaccharomyces pombe}		94312557 (3283, 3284) Novel Protein sim. GBank gil1575333 (U60416) - myr 6 myosin heavy chain [Rattus norvegicus]				95362691 (3291, 3292) Novel Protein sim. GBank gi 1076802 pir S49915 - extensin like protein - maize	94278428 (3293, 3294) Novel Protein sim. GBank gij5002573 emb CAB44338.1 - (Y17466) alpha-N-acetylgalactosamine alpha-2,6- sialyttransferase [Fugu rubripes]	
94143185 (3279, 3280)	87625160 (3281, 3282)	94312557 (3283, 3284)	1643 94131766 (3285, 3286)	88095125 (3287, 3288)	95013858 (3289, 3290)			1648 87642098 (3295, 3296)
1640	1641	1642	1643	1644	1645	1846	1647	1648

					CONTROL CONTROL POPULACION CONTROL
1649	95347628 (3297, 3298)	95347628 (3297, 3298) Novei Protein sım. (38ank gil834065)emb CAA38337 -	<u> </u>	Caunerin	204400, 222/0333, 33030200, 222/0330,
		(Sosalinani nerpesvilus of			20234922 20234924 20334825 20334826
					28551022, 28551024, 28551025, 28551025, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 100000, 100000, 100000, 100000, 100000, 100000, 100000, 100000, 100
					23331027, 23331020, 204303, 204307, 66743503 264008 52644045 264909
					56182435, 264511, 265007, 265008, 265009.
					264591, 264593, 60433438, 264596,
					55812038, 21906754, 265011, 264601,
					264602, 265017, 265018, 265019, 264682,
					264448, 264764, 264683, 264288, 264766,
					264685, 264687, 264768, 264688, 264769,
_					52644229, 264689, 21906765, 21906766,
_					21906767, 21906768, 55811957, 35695917,
					265021, 265022, 52644150, 264692,
					33657023, 33657109, 20281149, 18108370,
					264628, 18108374, 18108376, 35696423,
					35695855, 264632, 264634, 264635, 264636,
_					18108380, 264639, 264558, 18108382,
					18108384, 18108385, 18108387, 264080,
					264404, 60432113, 22279000, 22279002,
					264482, 264565, 264566, 264487
1650	87418539 (12999 1300)	87418539 (3209 3300) Novel Protein sim GRank nil3647335 Jemb JCAA210591.			265011, 264602, 21906767, 18108374,
		(AL031644) possible zinc-finger protein			18108377, 18108385
_		[Schizosaccharomyces pombe]		0004	254409 E254515E 18108307 35505285
[60L	91639773 (3301, 3302)	91639773 (3301, 3302) Novel Protein Sim. GBank gil4664270 empl. Ab43247.11		3911111636	204400, 25040100, 10100001; 00004001
		(AL050037) hypothetical protein [Homo sapiens]			222/0990, 222/0999, 204209, 2939/022,
_					29331824, 29331825, 29331826, 29331627,
					Z93318Z6, Z043U8, Z049U8, Z933183U,
	_				264910, 60432229, 21906754, 265010,
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					264288, 264688, 21906765, 21906766,
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					264693, 264629, 35695855, 264556, 264637.
					264557, 264559, 83373044, 56526486.
					22279000, 22279002, 264564
1652	86598622 (3303, 3304)	86598622 (3303, 3304) Novel Protein sim. GBank gi 1657837 (U73200) - p116Rip	Contains protein domain (PF00169) - struct	struct	22278997, 29146498, 56182435, 21906754,
		[Mus muscufus]	PH domain		264369, 21906765, 21906768, 21906769,
					265020, 52644150, 33657109, 22279000,
					22279002
1653	94255993 (3305, 3306)	94255993 (3305, 3306) Novel Protein sim. GBank gij3776054 jembjCAA06273 j	Contains protein domain (PF00047) - glycoprotein	glycoprotein	18108398, 22278995, 22278998, 264259,
		(AJ004999) Tapasin [Gallus gallus]	Immunoglobulin domain		29331822, 29331824, 60432289, 29331826,
					29331827, 29331830, 264909, 265006,
					265009, 60432229, 60433335, 60433436,
					21906754, 265017, 265019, 264448, 264683,
					264288, 265021, 265022, 264692, 16108364, 66374764 48408384 60432443 264667
7					22674751, 10100304, 00432113, 204301
1654	(18756471 (3307, 3308)	}		UNCLASSIFIED	133037 103, 204030

	Ω:	a_ma_bind 29331827, 265009, 21906766, 21906767, 265020, 265022, 33657109, 264638, 56526486, 264482		UNCLASSIFIED 52646317, 21906766, 21906767, 21906768, 87168518, 22278996, 265020, 22278999, 87168559, 264603, 265017, 264631, 265018, 265019, 22279002, 284482, 264635, 264565	UNCLASSIFIED 264488, 35696286, 264259, 35696052, 264508, 264509, 264905, 264906, 264907, 264908, 264909, 264510, 264511, 265007, 264909, 264510, 264511, 265006, 265007, 264596, 265010, 264600, 264602, 265017, 265018, 264600, 264602, 264020, 264089, 264766, 264766, 264769, 264789, 21906766, 35695917, 264690, 33657023, 264693, 33657109, 264639, 35696423, 35698655, 264634, 264635, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264637, 264637, 264639, 264637, 2
), S	UN	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00829) - UNCLASSIFIED Ribosomal prokaryotic L21 protein	Nn	<u>S</u>
1655 86689346 (3309, 3310) Novel Protein sim. GBank gij335577 emb CAA73496 - (Y13053) seryl-tRNA synthetase [Zea mays]	79962297 (3311, 3312) Novel Protein sim. GBank gi 1890141 db BAA18947 - (D83206) P24 protein [Mus musculus]	87771994 (3313, 3314) Novel Protein sim. GBank gif4557645frefiNP_001524.1[pHNRP - heterogeneous nuclear ribonucleoprotein L	87773778 (3315, 3316) Novel Protein sim. GBank gij3877072 emb CAA87060 - (Z46937) similarity with ribosomal protein L21 [Caenorhabditis elegans]	88230101 (3317, 3318) Novel Protein sim. GBank gij539218lpirj S38038 - hypothetical protein YKL201c - yeast (Saccharomyces cerevisiae)	94315313 (3319, 3320) Novel Protein sim. GBank gi[2497012[sp[Q10010]vSv4_CAEEL - HYPOTHETICAL 26.6 KD PROTEIN T19C3.4 IN CHROMOSOME III
86689346 (3309, 3310)	T				94315313 (3319, 3320)
165	1656	1657	1658	1659	1660

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### RBD, or RNP domain) #### RBD, or RNP domain) #### RBD, or RNP domain ###################################	<u> </u>			ONA monthism mater (Pruduto) -	UNCLASSIFIED	264488, 2227898, 264259, 29331824,
# ## Probain is min. CBank				AND ECOGNICION MON. (a.k.a. KKM,		29331826, 29331827, 29331828, 264509,
94135172 (3323, 3324) Novel Protein sim. GBank gilds 17EANSMEMBRANE protein sim. GBank gilds 4136[emb[CAB43275.1] - Contains protein domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WWV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - WAV domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] - kinase (AL050107) hypothetical pro				KBU, or KNP domain)		66712502, 29331830, 264908, 52644045,
94135172 (3323, 3324) Novel Protein sim. GBank gil 1736002is.ple2875iPzz_MOUSE - TRANSMEMBRANE PROTEIN PFT27 MOUSE - TRANSMEMBRANE PROTEIN PFT27 MOUSE - TRANSMEMBRANE PROTEIN PFT27 MOUSE - TRANSMEMBRANE PROTEIN PT27 MOUSE (3325, 3328) Novel Protein sim. GBank gil 3043692idbj BAA25510j - WWW domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] WWW domain (ABD11156) KIAA0564 protein film sapiens]						265007, 264512, 60433356, 60433438,
94135172 (3323, 3324) Novel Protein sim. GBanix gill 73060215pl22873FPZ7_MOUSE - TRANSMEMBRANE gill 73060215pl22873FPZ7_MOUSE - TRANSMEMBRANE gill 73060215pl22873 (ALGS0107) hypothetical protein [Homo sapiens] WW domain [ALGS0107) hypothetical protein [Homo sapiens] WW domain WW domain WW domain WM domain						55812038, 21906754, 265019, 264448,
94135172 (3323, 3324) Novel Protein sin. GBank gij7305021sppP227_MOUSE - TRANSMEMBRANE PROTEIN PF127 PROTEIN PF127 (AL 050107) hypothetical protein [Homo sapiens] (AL 050107) hypothetical protein [Homo sapiens] (AB011156) KIAA0584 protein [Homo sapiens]						264766, 264768, 264769, 21906768,
94135172 (3323, 3324) Novel Protein sim. GBank Proz. MOUSE - TRANSMEMBRANE PRIOTEIN PF127 Protein sim. GBank gil4884136jembjCAB43275.11 Contains protein domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] www.domain (AL050107) hypothetical protein [Homo sapiens] WWW.domain (AL050107) hypothetical protein [Homo sapiens] (AL05011156) KIAA0584 protein [Homo sapiens]					•	21906769, 265020, 33657023, 33657109,
94135172 (3323, 3324) Novel Protein sim. GBank gij4884136jembjCAB43275.1 - (AL050107) hypothetical protein [Homo sapiens] 94234076 (3327, 3328) Novel Protein sim. GBank gij3043682[dbjjBAA25510] - (AB011156) KIAA0584 protein [Homo sapiens] 94234076 (3327, 3328) Novel Protein sim. GBank gij3043682[dbjjBAA25510] -						65274791, 87168518, 264482, 264563,
94217146 (3325, 3328) Novel Protein sim. GBank gij8043692[dbj BAA25510] - WW domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens]						264564, 264565, 264567
94217146 (3325, 3326) Novel Protein sim. GBank gild884136jemblCAB43275.11 - Contains protein domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens] WW domain WM domain	1662	94135172 (3323, 3324)				18108392, 29331822, 29331828, 20281100,
94217146 (3325, 3326) Novel Protein sim. GBank gil3043892(bb) BAA25510 - Contains protein domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens]	_		gi 1/30502 sp P52875 PF27_MOUSE - TRANSMEMBRANE			264106, 265008, 265007, 265008, 18108348,
94217146 (3325, 3326) Novel Protein sim. GBank gild 884 136jemb[CAB43275.1] - WW domain (PF00397) - kinase (AL050107) hypothetical protein [Homo sapiens]	_		PROTEIN PFT27			21906766, 18108365, 18108366, 18108374,
9421/140 (3323, 3328) Novel Protein Sim. GBank gil3043692[dbjjBAA25510] - Kinase (AL050107) hypothetical protein [Homo sapiens] WW domain (PF00397) - Kinase (AL050107) hypothetical protein [Homo sapiens] WW domain (PF00397) - Kinase (AL050107) hypothetical protein [Homo sapiens] UNCLASSIFIED (AB011156) KIAA0584 protein [Homo sapiens]	30,					83373044, 18108385
(ALUSU107) hypothetical protein [Homo sapiens] WWW domain [94234076 (3327, 3328) Novel Protein sim. GBank gij3043892[dbj]BAA25510] - (ABD11156) KIAA0584 protein [Homo sapiens]	3		Novel Protein sim. GBank gi[4884136]emb[CAB43275.1] -	Contains protein domain (PF00397) -	kinase	52645156, 56182575, 22278994, 22278995,
84234078 (3327, 3328) Novel Protein sim. GBank gij3043892 dbj BAA25510] - (AB011156) KIAA0584 protein [Homo sapiens]			(ALUSU107) hypothetical protein [Homo sapiens]	WW domain		35696286, 22278996, 56994075, 22278997,
94234076 (3327, 3328) Novei Protein sim. GBank gij3043692 dbj BAA25510] - (AB011156) KiAA0584 protein [Homo sapiens]		_				22278998, 22278999, 264259, 29331822,
94234076 (3327, 3328) Novel Protein sim. GBank gij3043692 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						29331826, 29331827, 29331828, 33656970,
94234076 (3327, 3328) Novel Protein sim. GBank gij3043692 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						29331830, 264908, 56182435, 264511,
94234076 (3327, 3328) Novel Protein sim. GBank gij3043692 dbj BAAZ5510 - (AB011156) KIAA0584 protein [Homo sapiens]						60433356, 33657402, 33109954, 87168474,
94234076 (3327, 3328) Novel Protein sim. GBank gil3043892 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						87168559, 265017, 265018, 264605,
94234076 (3327, 3328) Novel Protein sim. GBank gil3043692 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo saplens]						18108351, 264764, 264288, 264766, 264768,
94234076 (3327, 3328) Novei Protein sim. GBank gij3043692/dbjjBAA25510j - (AB011156) KIAA0584 protein [Homo sapiens]						21906765, 21906766, 21906767, 21906768,
94234076 (3327, 3328) Novel Protein sim. GBank gi 3043692 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						21906769, 265021, 265022, 264691,
94234076 (3327, 3328) Novel Protein sim. GBank gij3043692 dbj BAA25510						33657023, 264693, 263967, 33657109.
94234076 (3327, 3328) Novel Protein sim. GBank gij3043692 dbj BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						264630, 52644332, 83373044, 87168518,
94234076 (3327, 3328) Novel Protein sim. GBank gi 3043692 db BAA25510 - (AB011156) KIAA0584 protein [Homo sapiens]						60432113, 22279000
·			Novel Protein sim. GBank gij3043692 dbj BAA25510 -		UNCLASSIFIED	264488, 263994, 35696286, 29331824,
264907, 264908, 264910, 264510, 26451 265009, 264910, 60170831, 264591, 264 264595, 87168474, 265011, 264600, 264 264504, 264605, 264760, 264760, 264760, 264762, 18108 2646081, 264605, 264760, 264762, 21906764, 21906764, 21906764, 21906764, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906764, 21906765, 21906			(AB011156) KIAA0584 protein [Homo sapiens]			35696052, 264508, 264509, 264905, 264906,
265009, 264910, 60170831, 264591, 264 264595, 87168474, 265011, 264600, 264 264595, 87168474, 265011, 264600, 264 264605, 264760, 264762, 18108 264681, 264682, 264763, 264762, 18108 264769, 21906764, 21906765, 219067						264907, 264908, 264909, 264510, 264511.
264595, B7168474, 265011, 264600, 264 264604, 264605, 264762, 18108 264604, 264605, 264762, 18108 264604, 264681, 264762, 18108 264681, 264882, 264763, 264683, 264763						265009, 264910, 60170831, 264591, 264592,
264604, 264605, 264760, 264762, 18108 264681, 264682, 264763, 219067763,						264595, 87168474, 265011, 264600, 264601,
264681, 264682, 264763, 264683, 26478 26428, 264684, 264766, 264676, 264684, 264766, 2906765, 21						264604, 264605, 264760, 264762, 18108351.
264288, 264684, 264766, 264687, 264766 21906765, 21906765, 21906765, 21906765, 21906765, 21906765 2190						264681, 264682, 264763, 264683, 264764,
264769, 21906764, 21906765						264288, 264684, 264766, 264687, 264768,
35695917, 265021, 264534, 60170615, 264690, 264691, 264692, 33657109. 33657182, 264628, 18108370, 264629, 35696423, 35695855, 264634, 264635, 264635, 264637, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264588, 33373044, 87168518, 264563, 264564, 87168518, 264563, 264564, 87168518, 264563, 264564, 87168518, 264563, 264564, 87168518, 264563, 264564, 87168518, 87168518	_					264769, 21906764, 21906765, 21906767,
264690, 264691, 264692, 33657109. 33657182, 284628, 18108370, 264629, 35696423, 35695855, 264634, 264635, 264635, 264634, 264638, 264636, 264634, 264638, 264636, 264636, 264638, 264638, 264638, 264638, 264638, 264589, 264589, 264589, 264589, 264589, 264589, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264569, 264586						35695917, 265021, 264534, 60170615,
33657182, 284628, 18108370, 264629, 35696423, 35695855, 264634, 264635, 264635, 264637, 264633, 264635, 264636, 264636, 264637, 264638, 264638, 264638, 264638, 264638, 264638, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264588, 264586						264690, 264691, 264692, 33657109,
35696423, 35695855, 264634, 264635, 264633, 264635, 264637, 264633, 264633, 264633, 264633, 264633, 264638, 264638, 264638, 264538, 264538, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264568, 264563, 264568, 264564, 264568, 264564, 264586						33657182, 264628, 18108370, 264629,
264555, 264636, 264637, 264638, 264638, 264638, 264563, 264559, 83373044, 87168518, 264563, 264563, 264563, 264563, 264563, 264569, 264586						35696423, 35695855, 264634, 264635,
264558, 83373044, 87168518, 264563, 264566, 264486						264555, 264636, 264637, 264638, 264639,
264566, 264486				-		264558, 83373044, 87168518, 264563,
						264566, 264486

	91226952 (3329, 3330)	1665 91226952 (3329, 3330) Novel Protein sim. GBank gi 1083506 pir S50065 -	Contains protein domain (PF00047) - immunoglob Immunoglobulin domain		264488, 29331826, 29331828, 264509. 264906, 264907, 264909, 264510, 264511. 264910, 264592, 264593, 264595, 264758. 264596, 264600, 264760, 264762, 264764, 264768, 264768, 264630, 264634. 264636, 83373044, 264564, 264566, 264567,
1666	95358160 (3331, 3332)	95358160 (3331, 3332) Novel Protein sim. GBank gil3913431 sp[O42643 DDXB_SCHPO - PUTATIVE PRE- MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE SPAC10F6.02C	Contains protein domain (PF00575) - helicase S1 RNA binding domain		26486 56994075, 22278999, 264259, 29331824, 29331826, 29331827, 29146498, 265009, 33109954, 87168559, 265019, 264288, 264688, 21806767, 21906769, 264691, 23557182, 18108370, 18108374, 18108385,
1667	91228655 (3333, 3334)	91228655 (3333, 3334) Novel Protein sim. GBank gij5689535 dbj BAA83051.1 - (AB029022) KIAA1099 protein [Homo sapiens]	Contains protein domain (PF01412) - transport Putative GTP-ase activating protein for Arf		264259, 29331822, 29331826, 264905, 264906, 264908, 264908, 264510, 265009, 264595, 264758, 265019, 265019, 265019, 265019, 265019, 265019, 26448, 264766, 264686, 21906765, 21906767, 21906769, 265020, 265021, 18108368, 18108370, 263972, 55810764, 264555, 83373044, 60432113, 22279000, 22279002
1668	88095135 (3335, 3336)	88095135 (3335, 3336) Novel Protein sim. GBank gi 2076894 gb AAB53983.1 - (AF002197) short region of weak similarity to protein kinase C; contains similarity to Pfam domain PF00130 (DAG_PE-bind), Score=10.0, E-value=0.0034. N=1 [Caenorhabditis elegans]		kinase	66714117, 284508, 264509, 264906, 264907, 264908, 264907, 264908, 264511, 264910, 264764, 264687, 264689, 33657109, 35696423, 35695855, 264632
1669	91227846 (3337, 3338) ,	91227846 (3337, 3338) Novel Protein sim. GBank gil3875371 lemb[CAA85414.1] - (Z36948) contains a valine and arginine rich domain, possesses weak similarity with the RNA binding domains from RNA splicing factor UZAF 65 KD subunit; cDNA EST EMBL:D64658 comes from this gene; cDNA EST EMBL:D66829 comes from		UNCLASSIFIED	29331825, 33109954, 264369, 264767, 264689, 33657109, 83373044
1670	87628009 (3339, 3340)			UNCLASSIFIED	264259, 29331824, 29331827, 60433438. 265022, 264636
1671	87346372 (3341, 3342 <u>)</u>	87346372 (3341, 3342) Novel Protein sim. GBank gil462451 splP34244 KKK1_YEAST - PROBABLE SERINE/THREONINE-PROTEIN KINASE YKL101W	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain		35696286, 22278997, 29331825, 264909, 21906754, 265017, 265018, 265019, 264682, 264683, 264768, 264688, 21906766, 21906767, 21906768, 21906769, 264556, 264556, 26279000, 264566
1672	86291834 (3343, 3344)	86291834 (3343, 3344) Novel Protein sim. GBank gi 1814270 (U74586) - double- stranded RNA specific adenosine deaminase [Rattus norvegicus]		deaminase	264906, 264909, 264632, 18108381

1673	88095137 (3345, 3346)	1673 88095137 (3345, 3346) Novel Protein sim. GBank gi 2076894 gb AAB53983.1 -		kinase	264488, 264569, 18108394, 56994075,
		(AF 002197) short region of weak similarity to protein kinase			22278996, 264259, 35696052, 264508,
		C; contains similarity to Pram domain PF00130 (DAG_PE-	domain (C1 domain)		264905, 264509, 264906, 264907, 264908,
		Diriul, Score=10.0, E-Value=0.0034, N=1 (Caenorhabdilis			264909, 264510, 264910, 60170831, 264592,
		elegans			264594, 264595, 264758, 264601, 264760,
					264762, 264683, 264764, 264288, 264766,
					264686, 264768, 264687, 264769, 264689,
					264690, 33657023, 264692, 264693,
					33657109, 264628, 264629, 18108374,
					35696423, 35695855, 264631, 264632,
					264634, 264635, 264637, 264556, 264638,
					264639, 264563, 264482, 264564, 264565,
į					264566, 264567, 264486
16/4		88238028 (3347, 3348) Novel Protein sim. GBank gi 5262467 emb CAB45693.1 -		kinase	29331822, 29331824, 264906, 52644045,
		(ALVOUGE) nypoinelical protein [Homo sapiens]			60433356, 87168559, 264448, 264288, 264686, 264691
1675		87606466 (3349, 3350) Novel Protein sim. GBank gij3128366 (AF010496) - 50S		UNCLASSIFIED	56181686, 35696286, 22278997, 22278998.
		ribosomal protein 19 [Rhodobacter capsulatus]			264259, 29331824, 29331827, 35696052,
					66712502, 264764, 264288, 264686, 264687,
					35695917, 265020, 264690, 264693,
					35695763, 18108370, 35696423, 35695855,
					264637, 264639, 18108385, 264564
9/91		95358086 (3351, 3352) Novel Protein sim. GBank gil4164065[gb]AAD05327 -		UNCLASSIFIED	264259, 29331827, 29331828, 264106,
		(Ar i i usi) latrophilin s spirce variant boat [Bos taurus]			264907, 265009, 264600, 265019, 264288,
					21906765, 265020, 265022, 35695855,
	11 100 07 07 07 07 07 07				83373044, 18108385
2		8/40858/ (3353, 3354) Novel Protein sim. GBank gij3327046 dbj BAA31591 - (AB014516) KIAA0616 protein [Homo sapiens]		UNCLASSIFIED	264908
1678				UNCLASSIFIED	29331824, 264102
1679		91214106 (3357, 3358) Novel Protein sim. GBank gil550452 (U08469) - 3-	Contains protein domain (PF00289) - carboxylase	carboxylase	264488, 18108392, 18108394, 52646842,
		methylcrotonyl-CoA carboxylase, biotin-carrier domain	Carbamoyl-phosphate synthase	-	18108397, 18108398, 35696286, 29331824,
		[Glycine max]	(CPSase)		265006, 265007, 265008, 265009, 18108348,
					265011, 18108351, 264683, 18108354,
					18108358, 18108359, 21906765, 29148627,
					29148629, 264690, 18108361, 18108362,
					18108364, 18108365, 18108368, 264628.
					18108379, 35696423, 35695855, 264635,
					18108381, 18108382, 18108383, 18108384,
1680	04006979 (2960 2960)	Manual Destain and Control of the Co			18108385, 18108388
0001	91003372 (3339, 3360)	91003372 (3339, 3300) Novel Protein Sim. GBank gip2394478 (AFU24500) - No		transport	65274572, 22278994, 22278999, 66714117,
		definition line tound [Caenornabditts elegans]			29331827, 55182435, 21906754, 265018,
1681	04324460 /3364 23631	0437450 (3269 2369) Manal Dankin in CBC 1215-12160 (2269 1369)		T	264288, 21906769
8	(2000, 1000) 001 +20+6	Novel Protein sim. Gbank gijobasos/jabjjbAA83052.1 -		UNCLASSIFIED	22278996, 29331822, 264908, 264593,
		(cyceyon)			264604, 263019, 264683, 33811937, 264630, 33657023, 35696423, 83373044, 264563
1682	86042710 (3363, 3364)			UNCLASSIFIED	264909, 265017, 264605
				1	

ś	3 94316213 (3365, 3366)	1683 94316213 (3365, 3366) Novel Protein sim. GBank	Contains protein domain (PF01852) - UNCLASSIFIED	UNCLASSIFIED	263994, 35696286, 35696052, 264508,
		gi 5031717 ref NP_005704.1 pGPBP - goodpasture antigen-	START domain	_	264509, 264905, 264906, 264907, 264908,
		binding protein		_	264909, 264511, 265006, 265007, 265009,
				_	264910, 264593, 264758, 265010, 265011,
					264760, 264761, 264762, 264763, 264764,
					264369, 264766, 264768, 35695917, 264692,
					33657109, 35696423, 35695855, 264634,
_					264635 264636 264638 264639 83373044
					264486
1684	4 80063409 (3367 3368)			LINCI ASSIFIED	264563 264566
1885	_	94323182 (3380 3370) Novel Protein sim CBank nil 1255371 (1153147) - Coded for	Contains protein domain (DE00625)	Linge	E0424170 52546842 22278004 35606286
<u>.</u>		hy C. elegans cDNA w/34a9 5: coded for by C. elegans			22278008 264260 52645080 20131824
		CONA vk34a9 3 Similar to quanylate kinase			24331826 265007 33657084 265018
		[Caenothabditis elegans]			264681 264448 264683 264369 264689
					21906765 21906767 21906768 21906769
					265021 264692 65274620 33657109
					2748555 264635 5264433 5618232
					22228000
1686	3 87820710 (3371, 3372)	87820710 (3371, 3372) Novel Protein sim. GBank gil2244707ldbjlBAA21115.11 -		UNCLASSIFIED	56182575, 264259, 264508, 264905, 264509,
		(AB005287) thrombospondin 1 [Bos taurus]	-		264907, 264908, 264510, 264511, 265006,
					264512, 265009, 264910, 264758, 265010.
	_				285011 284805 18108351 264764 264766
	_				400.011, 204.000, 101.000.01, 204.004, 204.000,
					18108357, 264768, 18108362, 264628,
					264630, 264631, 264634, 264635, 264637,
	\neg				264638, 264639, 264565, 264486, 264567
1687	_	94719400 (3373, 3374) Novel Protein sim. GBank		UNCLASSIFIED	35695917, 264906, 264907, 264908, 264510,
_		gi 4680679 gb AAD27729.1 AF13295 - (AF132954) CGI-20			265006, 265007, 264910, 264556, 18108381,
	_	protein [Homo sapiens]			18108383, 265011
1688	1			UNCLASSIFIED	264905, 264910, 264760, 264629, 264555
168	_	94325049 (3377, 3378) Novel Protein sim. GBank gil4240193 dbj BAA74875.1 -		UNCLASSIFIED	264569, 35696286, 22278999, 264508,
		(AB020659) KIAA0852 protein [Homo sapiens]			264908, 264909, 56182435, 33657402,
					60433438, 55812038, 265017, 265018,
					264448, 264764, 264288, 264686, 29148629,
					35695917, 265020, 265021, 263972,
	_				18108374, 65274791, 83373044, 264089
1690	_	83255346 (3379, 3380) Novel Protein sim. GBank gij3800736 (AF031572) - seven-		cadherin	264369, 21906766, 264692, 264639,
	_	pass transmembrane receptor precursor [Mus musculus]	Cadherin domain		87168518
1691		88095223 (3381, 3382) Novel Protein sim. GBank gi 2773208 (AF039713) - No			264768, 33657109, 29331827, 29148629,
		definition line found [Caenorhabditis elegans]			264510, 264106, 264910, 264109, 264508,
					60170831, 264563, 264905, 264564, 264691.
					264637, 264628, 264907, 264908, 33657023.
					264567, 264766, 263974
1692					264106
1693		87012775 (3385, 3386) Novel Protein sim. GBank	Contains protein domain (PF00042) - UNCLASSIFIED	UNCLASSIFIED	29331826, 264508, 264905, 264907, 264595,
_		gij121271 sp P02207 GLB_LAMFL - GLOBIN	Globin		265010, 265011, 21906768, 33657023,
					264629, 263978, 264558

3388)	IPOLR - polymerase (RNA) I (13.3kD)	Contains protein domain (PF01193) - mapolymerase RNA polymerases L / 13 to 16 kDa subunit	mapolymerase UNCLASSIFIED	35696286, 22278996, 22278999, 264259, 264259, 264259, 29331826, 29331826, 29331826, 29331826, 29331826, 29331828, 35696052, 2946499, 264905, 265008, 2644045, 264511, 265006, 265007, 265009, 265011, 18108351, 264763, 24682, 264448, 264683, 264683, 264693, 18108370, 18108374, 263978, 36596423, 35695855, 264556, 18108381, 18108385, 81168518, 264486, 264486
	87824038 (3391, 3392) Novel Protein sim. GBank gil4220517 emb CA422990 - (AL035356) hypothetical protein [Arabidopsis thaliana] 85740963 (3393, 3394) Novel Protein sim. GBank gil505652 (U10362) - GP36b		głycoprotein	. 22278997, 264259, 265010, 18108351, 264764, 21906766, 18108370 264682
1 -	87445285 (3395, 3396) Wycel Protein sim. Sepach gli (1995, 3396) Wycel Protein sim. GBank gli (1995, 3396) gli (1995, 3396) GBank gli (1995, 3396) axotrophin [Mus musculus]			56994075, 22276998, 22276999, 264509, 33657402, 264758, 87168474, 87168559, 265017, 265018, 26448, 264687, 29148627, 21906769, 29148629, 265020, 265022, 33657023, 264558, 87168518, 22279002
87424793 (3397, 3398)			UNCLASSIFIED	35696286, 264635
	87859161 (3399, 3400) Novel Protein sím. GBank gi 543344 pir S41647 - zinc finger 5 protein - mouse	Contains protein domain (PF00096) - nucl_recpt Zinc finger, C2H2 type	nucl_recpt	29331824, 52644045, 265008, 265009, 263969, 263971
86570488 (3401, 3402)			UNCLASSIFIED	264092, 264110, 263977
	87795092 (3403, 3404) Novel Protein sim. GBank gil3877439 emb CAA96652 - (272510) similarity to yeast UTR3 protein (Swiss Prot		МНС	22278995, 22278997, 264092, 29146498, 29146499, 264107, 264508, 264907, 264110,
	accession number P21374); cDNA EST EMBL:D72822 comes from this pene: cDNA EST FMBI: D75763 comes			264112, 265009, 60170831, 21906754, 265011, 265017, 264762, 1810826, 264288
	from this gene; cDNA EST yk274e3.3 comes from this			21906765, 35695917, 265021, 60170615,
	gene; cDNA EST yk274e3.5 c			263967, 33657109, 18108370, 263972, 263974, 18108374, 263976, 35695855,
				264555, 263981, 60170394, 18108385, 56526486, 87168518, 60432113
	79568651 (3405, 3406) Novel Protein sim. GBank gil451544 (U04267) - proline-rich cell wall protein [Gossypium barbadense]		UNCLASSIFIED	264909, 265017, 264628, 264629, 264638
_			UNCLASSIFIED	264369
	87795175 (3409, 3410) Novel Protein sim. GBank gil4519621(dbj BAA75670.1 - (AB017614) OASIS protein [Mus musculus]			264569, 35696286, 264907, 265010, 264687, 264768, 264692, 264693, 264636, 264566
				22278996, 22278998, 264259, 264509,
	gija rausajsplutatu i jauza_Human - HYPO i HETICAL PROTEIN KIAA0025		•	265018, 264764, 264685, 264686, 21906768, 21906768, 21906769, 265022, 264691, 264558,
]				7777

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18108396, 22278997, 264259, 29147620, 29331826, 29146498, 264905, 264906, 265008, 264593, 264595, 264758, 264596, 265018, 264760, 18108351, 264764, 264766, 264689, 264693, 18108370, 35698423, 264567	56182575, 22278996, 56994075, 264259, 29331822, 29331824, 29331826, 29331827, 29331828, 29146499, 29331830, 265009, 60170831, 33657402, 33109954, 87168559, 265019, 18108351, 264448, 21906765, 21906767, 21906768, 29148627, 29148629, 29148784, 60170615, 52644150, 33657023, 33657109, 18108374, 55811576, 264559, 18108385, 22279000, 264563	264107, 55811957, 263974, 263976, 263977, 263981	264556	65274572, 29331824, 29331826, 264768, 60431528, 35696423, 60432113, 264563	56994075, 264093, 264259, 29331822, 264099, 29331824, 29331827, 264107, 264110, 264511, 264592, 265011, 265018, 264683, 264686, 264689, 265020, 33657023, 263967, 33657109, 263974, 35696423, 35695855, 264630, 264636, 264566	22278995, 264508, 264758, 18108351, 18108370, 263974, 18108374, 264634, 56182323, 83373044, 60432113	22278997, 264757, 21906765, 265020, 265021, 264692, 56526486	264509, 264905, 264906, 264907, 264908. 264909, 264910, 264591, 265011, 264766, 264768, 264769, 264691, 264692, 264632, 284634, 264635, 264636, 264637, 264556, 264639, 264564	56182575, 29331824, 60432289, 264109, 264909, 265007, 264600, 265019, 264686, 265020, 2646893, 55811576, 264558, 60432113, 22279002
UNCLASSIFIED	synthase	UNCLASSIFIED	UNCLASSIFIED	sulfotransferase	kinase	UNCLASSIFIED		UNCLASSIFIED	transport
	Contains protein domain (PF01268) - synthase Formate-tetrahydrofolate ligase				Contains protein domain (PF00641) - kinase Zn-finger in Ran binding protein and others.	Contains protein domain (PF00643) - UNCLASSIFIED B-box zinc finger.			
1707 88041230 (3413, 3414) Novel Protein sim. GBank gil4321664 gb AAD15797 - (AF055470) ZNF258 [Homo sapiens]	91220519 (3415, 3416) Novel Protein sim. GBank gil5174591 [ref]NP_005947.1 pMTHF - 5.10- methylenetetrahydrofolate dehydrogenase, 5.10- methylenetetrahydrofolate cyclohydrolase, 10- formyltetrahydrofolate synthetase	0	[(91013729 (3421, 3422) Novel Protein sim. GBank gi 5031735 ref NP_005760.1 pHEC N-acetylglucosamine 6-O-sulfotransferase	95330184 (3423, 3424) Novel Protein sim. GBank gil5454168 ref NP_006453.1 pXAP4 - HBV associated factor	94143453 (3425, 3426) Novel Protein sim. GBank gi 160409 (M69183) - mature- parasite-Infected erythrocyte surface antigen [Plasmodium falciparum]		94260257 (3429, 3430) Novel Protein sim. GBank gi 5689537 db BAA83052.1 - (AB029023) KIAA1100 protein [Homo sapiens]	87400449 (3431, 3432) Novel Protein sim. GBank gil4589468 dbj BAA76761.1 - (AB012808) mBOCT [Mus musculus]
88041230 (3413, 3414)		80222583 (3417, 3418)	20754572 (3419, 3420)	91013729 (3421, 3422)			87420048 (3427, 3428)		
7	1708	1709		1711	1712	1713	1714	1715	1716

0/202220 (2423, 2424	1717 87553223 (3433, 3434) Novel Protein sim. GBank gi[2755411 emb CAA74749 -		UNCLASSIFIED	264569 264259 20331825 20331826
	(Y14391) GTP-binding protein [Homo sapiens]			29331828, 35696052, 264509, 264905,
				264907, 264908, 264909, 264512, 265009,
				264910, 264592, 264595, 264758, 264759,
				265017, 264681, 264764, 264766, 264686,
				18108357, 35695917, 264690, 264692,
				264693, 264628, 264629, 35696423, 264630,
,	•			264631, 264635, 264636, 18108380, 264638,
20,000000				264639, 18108388, 18108391
USZ628 (3435, 3436	6/U32628 (3435, 3436) Novel Protein sim. GBank		UNCLASSIFIED	285011, 264681, 264682, 264684, 264688,
	gi 2833262 sp Q14999 Y076_HUMAN - HYPOTHETICAL			264689, 21906765, 265021, 264691,
	PROTEIN KIAA0076 (HA0936)			33657023, 264693, 18108370, 35695855,
				264632, 264634, 264636, 18108388,
				22279002
1315259 (3437, 3438	94313259 (3437, 3438) Novel Protein sim. GBank		UNCLASSIFIED	18108396, 65274572, 35696286, 22278997,
	gi 4505197 ref NP_003473.1 pMLL2 - myeloid/lymphoid or			60432049, 56182181, 66714117, 60432289,
	mixed-lineage leukemia 2			29331826, 35696052, 29331828, 264906,
				29331830, 56182435, 264592, 60431735,
				60433438, 55812038, 264759, 265010,
				264600, 264601, 265017, 264448, 264764,
				264288, 264769, 21906766, 21906769,
				55811957, 265020, 265021, 52644150,
				33657023, 33657109, 33657182, 27486262,
				33657349, 35695763, 18108370, 60431528,
				18108374, 35696423, 55811576, 35695855,
		٠		264631, 56182323, 264559, 264564, 264486
4853063 (3439, 3440	94853063 (3439, 3440) Novel Protein sim. GBank gij2129478 pir S51939 -	<u> </u>	UNCLASSIFIED	56182575, 22278999, 264259, 29331824,
	chilinase (EC 3.2.1.14) precursor - beet	-		60432289, 29331827, 35696052, 264508,
				264905, 264906, 264907, 264908, 264909,
				264511, 264910, 264758, 21906754, 265011,
				264601, 264760, 264762, 264288, 264766,
				264686, 18108357, 264689, 21906765,
				55811957, 264693, 20281149, 264629,
				18108374, 55811576, 65274791, 264630,
				20281071, 264634, 264635, 264636, 264637,
				264556, 264638, 264639, 56182323,
799988 (3444 9449)	Months Brother also Charles also control to			87168518
91/22266 (3441, 3442)	Novel Protein sim. GBank gij4886461jemb CAB43381.1 -		UNCLASSIFIED	22278994, 22278999, 29331822, 265006.
	(Account inpolinence) profess (nomo sapiens)			265007, 265008, 55812038, 21906754,
				601/4639, 265011, 8/168559, 18108351,
				18108354, 21906765, 21906766, 21906768,
				21906/69, 265020, 3365/109, 18108370,
				18108374, 264556, 60170394, 83373044,
134549 (2443 3444)	94134549 (3443 3444) Naviel Protein cim CBank alise80275/JhilipA A 92000 11	100000000000000000000000000000000000000		18108385, 264486
ידידט (טדינט) טדינרטן	Appropriate Sim. Grank gijobba375 dajjBAA82968.1 -	Contains protein domain (PF00567) - kinase		56994075, 29331824, 29331828, 265009,
	(Addsub44) tudor repeat associator with PCTAIRE 2	Tudor domain	•	18108351, 21906768, 265020, 33657023,
	(Varius Holyegicus)			18108374, 83373044

264488, 264687, 264769, 21906767, 21906768, 56182575, 55811957, 22278997, 22278998, 265020, 264259, 264692, 33657023, 29331824, 29331824, 29331824, 29331825, 60432289, 33657182, 33656970, 33657349, 2946499, 264508, 264907, 18108370, 264629, 264501, 265007, 264908, 264909, 18108370, 264910, 264632, 264591, 60433229, 264595, 60433356, 264594, 60433438, 264595, 83373044, 55812038, 33109954, 33657084, 87168518, 87168474, 265010, 265011, 87168569, 264600, 60432113, 264604, 265019, 264764, 264488, 264389, 264766	264905, 18108359, 264693, 264628, 264631, 264636, 264555, 264556, 264558, 264559	35696286, 264259, 29331822, 35696052, 264508, 264509, 264905, 264906, 264907, 264908, 264909, 264900, 264909, 264910, 265009, 264591, 264601, 264760, 18108351, 264681, 264764, 264288, 264768, 294768, 21906769, 35695917, 264628, 35696423, 264630, 264631, 264632, 264635, 264638, 264648, 264648, 264648, 264648, 264648, 264648, 264648, 264648, 264648,	29146498, 264683, 264689	264905, 265011, 264689, 21906768
UNCLASSIFIED	UNCLASSIFIED	потеовох	UNCLASSIFIED	UNCLASSIFIED
•	gi 2340162 (AF005083) - dsRBP- Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF00023) - homeobox Ank repeat		
1723 95358181 (3445, 3446) Novel Protein sim. GBank gil4426962[gb AAD20633] - (AF126062) Arf-like 2 binding protein BART1 [Homo sapiens]	1724 87713806 (3447, 3448) Novel Protein sim. GBank gil2340162 (AF005083) - dsRBP- ZFa [Xenopus laevis]	1725 85655191 (3449, 3450) Novel Protein sim. GBank gij3152662 (AF064604) - KE03 protein [Homo sapiens]	(2)	85296362 (3453, 3454) Novel Protein sim. GBank 9146893481gb AAD27861.1 AF13256 - (AF132562) BCDNA, LD14270 [Drosophita melanogaster]
95358181 (3445, 3446	87713806 (3447, 3448	85655191 (3449, 3450	85754255 (3451, 3452)	, , , , , , , , , , , , , , , , , , ,
<u>27</u>	1724	1725	1726	1727

1728	95349515 (3455, 3456)	1728 [95349515 (3455, 3456) Novel Protein sim. GBank gil4406549 gb AAD20027 -		UNCLASSIFIED	60424179, 18108397, 56182575, 22278995,
		(AF131738) Unknown [Homo sapiens]			56994075, 35696286, 22278997, 22278998,
					22278999, 264094, 60432049, 264259,
					29331822, 29331824, 56182181, 29331825,
					60432289, 29331826, 29331827, 35696052,
			-		264905, 264906, 264907, 29331830,
					66712502, 264908, 56182435, 264511,
					265008, 265009, 60432229, 60433356,
					33657402, 60433438, 264759, 21906754,
					87168474, 265010, 265011, 87168559,
					265017, 265018, 265019, 55811150, 264681.
					264448, 264682, 264763, 264683, 264288,
					264684, 264369, 264685, 264766, 264687,
					264769, 21906764, 264689, 21906765,
					21906766, 21906767, 21906768, 35695917.
					265020, 265021, 265022, 264535, 264691,
					264692, 33657023, 264693, 33657109,
	_				18108370, 264628, 263972, 264629,
					18108374, 18108376, 55810764, 65274791,
					35695855, 264631, 264634, 264635,
					60431850, 264636, 264638, 60170394,
					264639, 83373044, 56526486, 87168518,
					60432113, 22279000, 22279002, 264564,
					264566
1729	91227948 (3457 3458)	91227948 (3457, 3458) Novel Protein sim, GBank gil854065 emb CAA58337 -		UNCLASSIFIED	264906, 264907, 264908, 264511, 264555.
2	()	(X83413) U88 (Human heroesvirus 6)			83373044, 264596, 264566
1730	85483474 (3459, 3460)			UNCLASSIFIED	29331822, 29331825, 29331828, 264907,
<u>:</u>	, , , , , , , , , , , , , , , , , , , ,				264908, 264909, 265011, 264764, 264629
1731	88266068 (3461, 3462)	88266068 (3461, 3462) Novel Protein sim, GBank gil631600 pir 1847094 -		UNCLASSIFIED	52646842, 264907, 264909, 56182435,
					55811386, 87168559, 265018, 265019.
					264760, 52644229, 55811576
1732	_	91218878 (3463, 3464) Novel Protein sim, GBank gil4240231 dbj BAA74894.1 -		struct	56182575, 29331822, 29331824, 29331827,
!		(AB020678) KIAA0871 protein [Homo sapiens]			66712502, 264591, 33657402, 60433356,
					265019, 21906768, 21906769, 35695917,
					265020, 265021, 264636, 56182323
1733	$\overline{}$	87617178 (3465, 3466) Novel Protein sim. GBank gil 1575756 (U70674) - m-Numb	Contains protein domain (PF00640) - synthase	synthase	264907, 264910, 33657402, 265010, 264681,
) :			Phosphotyrosine interaction domain		264683, 264684, 264686, 264769, 264691,
			(PTB/PID).		264692, 264693, 264628, 264636, 264556
1734	1734 87795261 (3467, 3468)				264693

264488, 18108394, 18108398, 22278998, 60432049, 264259, 29331822, 264908, 265006, 265007, 265008, 265009, 264591, 33657402, 265010, 265011, 87168559, 264769, 264689, 21906765, 21906768, 2914862, 2914874, 265022, 5244150, 18108364, 18108385, 33657109, 18108370, 18108380, 18108385, 87168518, 264563, 18108380	264490, 264259, 66714117, 66712502, 56182435, 265006, 265008, 264910, 60433356, 87166559, 265017, 265019, 18108351, 265020, 265022, 33657023, 33657109, 284555, 60431850, 264637, 60170394, 264558, 264639, 264564	29331822, 29331830, 264591, 265011, 265018, 265019, 265019, 22279002 264908, 264909, 265008, 264910, 264566	35696052, 264603, 264557	264604, 21906764, 18108364, 264629, 35695855, 264636 60432289, 29331627, 264509, 265009, 60432229, 264759, 265017, 264767, 264688, 264689, 21906769, 265020, 33657109	264906, 264910, 264758, 265011, 264631, 264638, 264566 265017, 265020, 264692	22278998, 264508, 264907	65274791, 264639, 264559	52646842, 29331824, 29331825, 52644045, 56182435, 265007, 52846317, 87168474, 265018, 18108351, 264369, 264769, 264689, 35695917, 60170615, 264691, 33657023, 65274791, 35695855, 60170394, 22279000, 264482
	UNCLASSIFIED	kinase UNCLASSIFIED	UNCLASSIFIED	traffic UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transcriptfactor
Contains protein domain (PF01169) - Uncharacterized protein family UPF0016		Contains protein domain (PF00168) - Kinase C2 domain UNCLA	Contains protein domain (PF00441) - UNCLASSIFIED Acyl-CoA dehydrogenase	traffic Contains protein domain (PF01363) - UNCLASSIFIED FYVE zinc finger			Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat	
1735 88318638 (3469, 3470) Novel Protein sim. GBank gil4836807[gb]AAD30566.1 AF14679 - (AF146793) PFT27 [Mus musculus]		88165549 (3473, 3474) Novel Protein sim. GBank gil2143607 pir S68695 - B/K protein - rat protein - rat protein sim. GBank gil2225941 emb CAA69714 -		83592939 (3479, 3480) Novel Protein sim. GBank gil4809[emb[CAA44309] - (X62452) YCR601 [Saccharomyces cerevisiae] (S6010100 (3481, 3482) Novel Protein sim. GBank gil4883898[gb]AAD31695.1]AF13042 - (AF130420) serine protease-like protein isoform [Homo sapiens]	85788814 (3483, 3484) Novel Protein sim. GBank gi 4505193 ref NP_003667.1 pMLD - membrane fatty acid (lipid) desaturase 86966475 (3485, 3486)	1744 91224003 (3487, 3488) Novel Protein sim. GBank gij728832 sp P39189 ALU2_HUMAN - III! ALU SUBFAMILY SB WARNING ENTRY III!	94326110 (3491, 3492) Novel Protein sim. GBank 94326110 (3491, 3492) Novel Protein sim. GBank 913131756 sp p38873 YHY6_YEAST - HYPOTHETICAL 175,8 KD PROTEIN IN GND1-IKI1 INTERGENIC REGION	94324333 (3493, 3494) Novel Protein sim. GBank gil 1658503 (U75467) - Atu Drosophila melanogaster

1748	88003580 (3495, 3496)	1748 88003580 (3495, 3498) Novel Protein sim. GBank gil4504511[ref]NP_001530.1[pHSJ2 - heat shock protein, DNAJ-like 2	Contains protein domain (PF00684) - eph OnaJ central domain (4 repeats)	eph	264489, 56182575, 29331824, 56182435, 264112, 265007, 265019, 264764, 21906768, 265020, 264681, 55811576, 264635, 264555
1749	83363091 (3497, 3498)	83363091 (3497, 3498) Novel Protein sim. GBank gij5650780jgbJAAD45948.1[AF15196 - (AF151968) RGS protein RGS-17 IGallus aallusl	Contains protein domain (PF00615) - oncogene Regulator of G protein signaling domain	oncogene	26456, 264557, 264559 264106
1750	94321664 (3499, 3500)	94321664 (3499, 3500) Novel Protein sim. GBank gjik996894 gb AAC2844.2 - (AF065164) hyperpolarization-activated, cyclic nucleotide- igated channel 2 [Homo sapiens]			33657402, 264288, 52644150, 263974, 83373044
1751	83373058 (3501, 3502)	83373058 (3501, 3502) Novel Protein sim. GBank gil2760161[db] BAA24184 - (AB010054) outer arm dynein light chain 2 [Anthocidaris crassispina]	Contains protein domain (PF00560) - ATPase_associated 265010, 264369 Leucine Rich Repeat	ATPase_associated	265010, 264369
1752	86456530 (3503, 3504)	86456530 (3503, 3504) Novel Protein sim. GBank gi 3915482 sp P74346 YG29_SYNY3 - HYPOTHETICAL 36.0 KD PROTEIN SLR1629	Contains protein domain (PF00849) - deaminase RNA pseudouridylate synthase	deaminase	264510, 264593, 264682, 21906765, 18108370
1753	94235159 (3505, 3506)	94235159 (3505, 3506) Novel Protein sim. GBank gi 2852636 (AF007155) - unknown [Homo sapiens]	Contains protein domain (PF01553) - phosphatase Acyltransferase	phosphatase	56994075, 22278996, 264908, 60170831, 264682, 264764, 264369, 264288, 264685, 264687, 21906766, 264692, 264693, 65274620, 65274791, 35695855, 264637, 264564
	88095323 (3507, 3508)		·	transport	264488, 35696286, 264509, 264906, 264907, 264908, 264909, 264511, 264910, 264591, 33657402, 264594, 264757, 264758, 264600, 264604, 264762, 264763, 264685, 264634, 264631, 264638, 264634, 264638, 264638, 264639, 264639, 264564, 264565, 264565, 264567
1755	ļ	79470282 (3509, 3510) Novel Protein sim. GBank gij1176422 (U43194) - rhophilin [Mus musculus]		UNCLASSIFIED	264686
1756		92962614 (3511, 3512) Novel Protein sim. GBank gil4432860 gb AAD20708 - (AC006300) putative glucose-induced repressor protein [Arabidopsis thaliana]			52646842, 22278994, 22278995, 56994075, 22278996, 22278997, 264259, 29331822, 60432289, 29331822, 60432289, 29331822, 265009, 60432289, 6043328, 60433438, 33109954, 21906754, 265017, 265018, 265021, 265018, 265021, 265018, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906767, 265022, 264822, 27486262, 27486264, 18108388, 81168518, 60432113, 22279000, 262799002, 26482
1757	85357380 (3513, 3514)	95357380 (3513, 3514) Novel Protein sim. GBank gi[5441615 emb CAB46856.1 - (AJ386557) zinc finger protein [Canis familiaris]	Contains protein domain (PF00096) - dna_rna_bind Zinc finger, C2H2 type	dna_rna_bind	22278997, 264259, 60432289, 29331827, 264908, 52640465, 265008, 264593, 265019, 264766, 21906768, 65274620, 18108385, 60432113, 264566, 264487

22278994, 22278999, 29331822, 29331824, 29331825, 33656970, 264508, 265006, 265007, 265009, 264591, 33657402, 33109954, 87168474, 264600, 265017, 265018, 21906769, 265020, 265021, 33657023, 33657109, 264629, 18108374, 264563	264759			22278999, 264259, 264905, 264907, 60170831, 265010, 265011, 265017, 26448, 21906765, 21906766, 21906767, 21906768, 265021, 264690, 33657109, 18108374, 264558, 60170394	56181686, 29331825, 35696052, 264905, 264906, 264906, 264909, 264763, 264682, 264769, 35695917, 265022, 33657023, 18108374, 35696423, 264634	22278998, 264259, 35696052, 29331828, 56182435, 265008, 265017, 265018, 264448, 264288, 21906766, 21906767, 29148627, 35695917, 264691, 33657023, 60432113, 22279002	264488, 18108394, 22278995, 22278997, 22278999, 264259, 264104, 264508, 264905, 22278999, 264259, 264104, 264508, 264905, 265009, 265009, 21906754, 265010, 265017, 264603, 265018, 265019, 18108351, 264682, 264369, 264288, 264766, 18108359, 21906766, 21906767, 29148627, 29148629, 35695917, 265021, 264639, 264628, 264629, 18108374, 263976, 264636, 83373044, 22279000, 22279002
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	desaturase		collagen	UNCLASSIFIED
·		·	Contains protein domain (PF00618) - Guanine nucleotide exchange factor for Ras-like GTPases; N-terminal motif	Contains protein domain (PF00173) - desaturase Heme-binding domain in cytochrome b5 and oxidoreductases		Contains protein domain (PF01410) - Fibrillar collagen C-terminal domain	
1758 87612971 (3515, 3516) Novel Protein sim. GBank gij3881040jembjCAA16403j - (AL021497) predicted using Genefinder [Caenorhabditis elegans]		87329716 (3519, 3520) Novel Protein sim. GBank gij5262748 emb CAB45688.1 - (AJ133120) Protine rich synapse associated protein 2 [Rattus norvegicus]	87409586 (3521, 3522) Novel Protein sim. GBank gil127749 sp P10569 MYSC_ACACA - MYOSIN IC HEAVY CHAIN	95319887 (3523, 3524) Novel Protein sim. GBank gil3169158 (AC004770) - BC269730_2 [Homo sapiens]	91224013 (3525, 3526) Novel Protein sim. GBank gil4809026lgb AAD30062.1 - (AF132856) suppressor of G2 allele of skp1 homolog [Homo sapiens]	87757697 (3527, 3528) Novel Protein sim. GBank gi 1360669 pir CGHU1V -collagen alpha 1(V) chain precursor - human	91230091 (3529, 3530) Novel Protein sim. GBank gil488806 pir S35503 - finger protein neuralized - fruit fly (Drosophila melanogaster)
87612971 (3515, 3516)	36994372 (3517, 3518)				9,1224013 (3525, 3526)		
1758	1759	1760	1761	1762	1763 -	1764	1765

62274572, 56182575, 56994075, 35696286, 22278997, 29331822, 29331828, 60432289, 29331828, 35696052, 29331830, 66712502, 29331828, 56182435, 264511, 265007, 265009, 60170831, 60433229, 60433438, 55812038, 21906754, 85658542, 87188559, 264601, 265017, 265018, 265019, 264762, 21906766, 21906766, 21906766, 21906767, 21906768, 265020, 265021, 265022, 264691, 33657109, 27486261, 33657349, 18108370, 18108372, 264634, 265638, 264639, 5618232, 264632, 264638, 264639, 56182323, 22279000, 22279000, 22279002, 264482, 264486, 264486, 264482, 264486, 264482, 264488,	264910	56994075, 29331826, 265008, 87168474, 265017, 265018, 264761, 55811150, 264764, 56181562, 264689, 21906765, 21906768, 21906769, 35695917, 264680, 33657023, 35695763, 60431528, 35696423, 55811576, 35695855, 22279000, 22279002, 264564	29331826, 29331827, 35696052, 264512, 265007, 265009, 265017, 265019, 264762, 18108351, 264769, 21906765, 21906766, 21906767, 21906767, 21906769, 265020, 265021, 264691, 264693, 18108370, 264556, 83373044	29148627, 35696286, 29147620, 265006, 265007, 265007, 265008, 18108385, 65274727, 264482, 264369, 264766	264107, 33657109, 56526486	264508, 264906, 264639	264259, 29331822, 264508, 264905, 264905, 264907, 264907, 264908, 265007, 265009, 264910, 264591, 264758, 264764, 264288, 264768, 264769, 264563	264768
UNCLASSIFIED		nuci_recpt	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		interferon
-					Contains protein domain (PF00807) - UNCLASSIFIED Apidaecin			
1775 95359330 (3549, 3550) Novel Protein sim. GBank gil1469199lbJBA09487 - (D50928) The KIAA0138 gene product is novel. [Homo sapiens]	94133756 (3551, 3552) Novel Protein sim. GBank gil4589676ldbjlBAA76857.1 - (AB023230) KIAA1013 protein [Homo sapiens]	87447171 (3553, 3554) Novel Protein sim. GBank gij3219939 sp P87115 YDK9_SCHPO - HYPOTHETICAL 116.5 KD PROTEIN C20G8.09C IN CHROMOSOME I	94851624 (3555, 3556) Novel Protein sim. GBank gij3875648 emb CAA91454.1 - (Z66561) Similarity to Human rab13 protein (PIR Acc. No. A49647). Contains the ATP/GTP-binding site motif (PROSITE PS00017); cDNA EST EMBL:M89412 comes from this gene; cDNA EST yk212g9.3 comes from this gene; cDNA EST yk212g9	94133758 (3557, 3558) Novel Protein sim. GBank gil4589676 db BAA76857.1 - (AB023230) KIAA1013 protein [Homo sapiens]			88094607 (3563, 3564) Novel Protein sim. GBank gi 729225 sp P41237 CTXN_RAT	85717905 (3565, 3566) Novel Protein sim. GBank gi 2257543 db BAA21436 - (AB004538) protein arginine N-methyltransferase Schizosaccharomyces pombe]
5 95359330 (3549, 3550)	94133756 (3551, 3552)	87447171 (3553, 3554)						
721	1776	1777	1778	1779	1780	1781	1782	1783

35696286, 264259, 35696052, 264508, 264905, 264906, 264907, 66712502, 264908, 264907, 66712502, 264908, 264909, 265007, 265008, 265009, 264910, 264591, 264511, 264617, 264602, 264604, 26405, 264762, 264763, 264763, 264763, 264763, 264697, 264567, 264486	65274572, 264259, 29331822, 29331824, 29331825, 29331825, 29331826, 264905, 264906, 264908, 66712502, 56182435, 264511, 265007, 60433356, 55811150, 264683, 264687, 52644229, 21906767, 52644150, 33657023, 65274620, 33657182, 65274791, 35695855, 264555, 65274727, 22279002	264908, 35696423, 264636	264488, 264905, 264908, 264909, 264595, 264764, 264766, 264692, 60431528, 264629, 264636, 264564, 264566	264488, 83373044	264488, 29331828, 264909, 18108351, 264288, 265021, 264555, 264636	35696052, 264905, 264906, 264907, 264908, 264909, 265008, 264910, 264758, 265011, 265019, 264764, 264766, 264769, 264628, 264635
struct	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	
Contains protein domain (PF00169) - struct PH domain						
95197093 (3567, 3568) Novel Protein sim. GBank gil1755049 (U55042) - myosin X [Bos taurus]	1785 95357475 (3569, 3570) Novel Protein sim. GBank gil4589552 dbj BAA76798.1 - (AB023171) KIAA0954 protein [Homo sapiens]	85296465 (3571, 3572) Novel Protein sim. GBank gi[17788 sp P26770 CYA4_RAT ADENYLATE CYCLASE, TYPE IV (ATP PYROPHOSPHATE-LYASE) (ADENYLYL CYCLASE)	87434784 (3573, 3574) Novel Protein sim. GBank gij3877175 emb CAA90338.1 - (Z50028) cDNA EST yk321h8.5 comes from this gene; cDNA EST EMBL:D68896 comes from this gene; cDNA EST yk395f9.5 comes from this gene [Caenorhabditis elecans]		88094529 (3577, 3578) Novel Protein sim. GBank gil2088669 (AF003130) - F55A12.9 gene product [Caenorhabditis elegans]	
95197093 (3567, 3568)	95357475 (3569, 3570)	85296465 (3571, 3572)	87434784 (3573, 3574)	91228779 (3575, 3576)		82489734 (3579, 3580)
1784	1785	1786	1787	1788		1790

264488, 264686, 264687, 264768, 18108394, 264769, 18108397, 264259, 264691, 264692, 33657023, 264693, 264699, 264692, 33657023, 264693, 264699, 264696, 264696, 264696, 264696, 264606, 264631, 265009, 264631, 264631, 264632, 264632, 264634, 264635, 264636, 264636, 264694, 264760, 264694, 264760, 264694, 264763, 264684, 264763, 264684, 264763, 264684, 264768, 264664, 264664, 264664, 264665, 264766, 264664, 264665, 264766, 264664, 264668, 264768, 264684, 264768, 264681, 264769, 264684, 264768, 264684, 264768, 264766, 264766, 264766, 264664, 264685, 264766	22278997, 264259, 264508, 265007, 33657402, 87168559, 264369, 33657023, 33695855, 20281071, 264559, 18108387, 87168518	Contains protein domain (PF01532) - ATPase_associated 6574572, 22278995, 22278996, 22278997, Glycosyl hydrolase family 47 Glycosyl hydrolase family 47 66714117, 60432289, 29331824, 264910, 265009, 60170831, 6043336, 21906754, 265010, 265010, 265017, 265010, 265017, 265010, 264910, 265010, 265017, 265010, 264910, 265010, 265017, 265010, 264910, 265010, 265010, 265010, 264910, 265010, 265010, 265010, 265010, 265020, 265020, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265021, 265022, 60170615, 52644150, 33657023, 33657109, 18108370, 18108374, 65274791, 20281071, 60432113, 22279000, 264482, 264564	18108394, 22278995, 22278999, 264259, 29331822, 29331824, 29331825, 29146498, 29146499, 264508, 264905, 52644045, 264712, 265006, 265908, 264910, 60433356, 264757, 55812038, 87168474, 265011, 265017, 18108351, 264763, 264448, 264683, 264359, 21906765, 21906766, 21906766, 21906765, 21906765, 33657023, 264629, 18108374, 18108376, 35696423, 35695855, 264556, 264557, 264638, 264558, 18108376, 264638, 264558, 18108376, 264638, 264558, 18108376, 264638, 264558, 18108376, 264638, 264558, 264556, 264567	264632, 264635, 264636, 264595, 264596, 264907, 284566, 264909	264488, 264907, 264909, 264594, 264595, 264766, 264687, 21906765, 21906767, 264628, 284630, 264559
glycoprotein	UNCLASSIFIED	ATPase_associale	UNCLASSIFIED	UNCLASSIFIED	glycopratein
Contains protein domain (PF00047) - glycoprotein Immunoglobulin domain	Contains protein domain (PF01585) - UNCLASSIFIED G-patch domain		Contains protein domain (PF01798) - UNCLASSIFIED Putative snoRNA binding domain		
1781 95197259 (3581, 3582) Novel Protein sim. GBank gi 2114321 db BAA20037 - (D88733) membrane glycoprotein [Equine herpesvirus 1]	87792690 (3583, 3584) Novel Protein sim. GBank gil4337106jgb AAD18082 - (AF129756) BAT4 [Homo sapiens]	95337877 (3585, 3586) Novel Protein sim. GBank gil5579331[gb]AAD45504.1 AF14573 - (AF145732) endoplasmic reticulum alpha-mannosidase I [Homo sapiens]	87759806 (3587, 3588) Novel Protein sim. GBank gil4914604 emb CAB43677.1 - (AL050369) hypothetical protein [Homo sapiens]		86599486 (3591, 3592) Novel Protein sim. GBank gijs85084fsplQ07803 EFGM_RAT - ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR (MEF-G)
95197259 (3581, 3582)	87792690 (3583, 3584)	95337877 (3585, 3586)		79747856 (3589, 3590)	
1791	1792	1793	1784	1795	1796

707	04223340 /2503 2504	04222240 (2502 2504) Namel Deskin Jim Charle Handonski (Handonski Handonski)		
:		[Arabidopsis thatiana]	ioosomaipioi	22210390, 22210391, 22210393, 22210393. 29331822, 264910, 60170831, 21906754.
				52644229, 21906765, 21906768, 21906769.
				35695917, 265022, 52644150, 264691,
1	_			33657023, 263967, 33657109, 22279000
8		91221270 (3595), 3599) Novel Fridein Sim. Chank git/832906[db] BAA24668.1	peptidase	22278994, 56994075, 22278997, 22278998,
		(Cososto) orbebuoyi pepuoase III [Kaiius noivegicus]		22278999, 264259, 29331826, 60432289,
				29331828, 33656970, 265008, 60432229,
				264757, 60433438, 21906754, 33657084,
				87168559, 265017, 18108351, 264682,
				264448, 264288, 21906765, 21906766,
				21906767, 21906768, 21906769, 35695917,
				265020, 265021, 33657023, 33657182,
				27486261, 27486265, 33657349, 263973,
				18108374, 55811576, 35695855, 18108385,
				87168518, 22279000, 264486
1799		86321713 (3597, 3598) Novel Protein sim. GBank gil5689541 dbj BAA83054.1 I -	ebh	264908, 21906754, 21906767, 21906769,
		[(AB029025) KIAA1102 protein [Homo sapiens]		265020, 33657023, 264692, 264693, 264404,
				22279000
8	1800 87080116 (3599, 3600)		UNCLASSIFIED	264691, 264556, 264566
<u>6</u>	95060723 (3601, 3602	95060723 (3601, 3602) Novel Protein sim. GBank		52644045, 265007, 264632
		gi 4680679 gb AAD27729.1 AF13295 - (AF132954) CGI-20		
		protein [Homo sapiens]		
1802		87771012 (3603, 3604) Novel Protein sim. GBank		35696286, 66714117, 264508, 264509,
		gil134920[sp P21997 SSGP_VOLCA - SULFATED		56182435, 264512, 18108351, 264688,
		SURFACE GLYCOPROTEIN 185 (SSG 185)		55811957, 264692, 55811576, 35695855,
5	2000, 20000000			264486
1803	95060725 (3605, 3606	95060725 (3605, 3606) Novel Protein sim, GBank		264686, 264488, 264687, 264489, 264768,
		gi 4680679 gb AAD27729.1 AF13295 - (AF132954) CGI-20		264769, 264689, 21906769, 35696236,
		protein [Homo sapiens]		35695917, 264259, 264691, 264692, 264693,
				20281099, 18108364, 35696052, 264508,
				264509, 264905, 264906, 18108370, 264628,
				264907, 66712502, 264908, 264909,
				18108374, 18108376, 35696423, 35695855,
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				265008, 264910, 264631, 264632, 264634,
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				264758, 264596, 18108385, 18108387,
				265011, 264760, 264563, 18108351, 264762,
				264564, 264448, 264565, 264763, 264683,
				264764, 264566, 264288, 264486, 264567.
1				264765, 264766

1804	אחפר לחפרו בחכחלללפן	1804 187770201 (1807 1808) Novel Protein sim GBank pil/1879914 lemblCAA98538 11.			52646365, 22278997, 22278999, 264905,
}	(222 (22) (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (222 (22) (222 (222 (222 (222 (222 (222 (22) (222 (22) (222 (222 (222 (22) (222 (22) (222 (222 (222 (222 (22) (222 (22) (222 (222 (22) (222 (22) (222 (222 (22) (222 (22) (222 (222 (22) (222 (22) (222 (22) (222 (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22) (222 (22))))))))				264908, 264909, 264910, 21906754, 264766,
		EMBI :C13850 comes from this cene: cDNA EST			21906765, 21906768, 35695917, 265020,
		EMBI :C11575 comes from this gene: cDNA EST vk343f4.5			265022, 264691, 264637, 264639, 22279000,
		comes from this gene [Caenorhabditis elegans]			264564, 264566
1805	95330375 (3609, 3610)	95330375 (3609, 3610) Novel Protein sim. GBank			29331824, 29331825, 29331826, 29331827,
		gil5453644trefINP 006461.1lpEBBP - estrogen-responsive			29331828, 87168559, 264288, 264687,
					52644229, 35696423, 264636, 60432113
1806	-	94133762 (3611, 3612) Novel Protein sim. GBank gil4589676 dbj BAA76857.1] -	S	struct	264094, 264105, 264908, 35696423, 265006,
	_				265007, 265008, 264555, 264592, 265011,
					265018, 264369
1807	86943032 (3613, 3614)			5	29331824, 264908, 264910, 33657023,
					263978
1808	_	87642711 (3615, 3616) Novel Protein sim. GBank gil4884079 emb CAB43235.1 -	2	UNCLASSIFIED	264488, 35696286, 66714117, 35696052,
					66712502, 264592, 60433438, 52644296,
_					265010, 264683, 264369, 264689, 55811957,
_					35695917, 33657109, 35695763, 55810764.
					18108379, 35696423, 35695855, 56182323,
	-				264563, 264564, 264487
400		Sevilation (2817 2817) (1979) Alaria cia cia and Charles (1979)		INCLASSIFIED	264594 55811150 264686 29148629.
80		postato (Mus musculus)			29148784 264690 264629 18108374.
					264556, 264557, 264558
4010		09006346 (2610 2620) Mayol Dratein clm CBank		INC! ASSIFIED	264488 35696052 264905 264906 264907,
2		nitasco44(sniP47179(V.)9P YEAST - HYPOTHETICAL			264908, 264909, 264511, 265009, 264910,
		1449 A KD BOOTEIN IN BATS DAI & INTERCENIC PECION			264592 264593 264594 33657402, 264757,
_		DECLIBEOD			264595 264758 264596 264759 264600.
		אספאסטשער			264601 264762 264683 264764 264288
					204001, 204702, 204003, 204704, 204200, 264684 264766 264767 264686 264768
					264887 264780 264020 264021 264690
					264601 264603 264628 264629 18108374
					264630 264631 264632 264634 264635
					264636, 264637, 264638, 264639, 264563.
					26456 26448 264567
1811	_	98086272 (3621 3622) Novel Protein sim (38ank oil) 134984 Injrill 37275 - death-	Contains protein domain (PF00023) - Ikinase	kinase	264488, 264259, 264508, 264509, 264905,
<u>.</u>		associated protein kinase (EC 2.7.1.) - human	Ank repeat		264906, 264907, 56182435, 264511, 264512.
					264910, 264758, 265011, 264600, 264604,
_					18108354, 264766, 264686, 264769, 264534.
					60170615, 33657023, 264629, 264631,
					264639, 264563, 264482, 264483
1812	79245772 (3623, 3624)				29331822, 29331824, 265019, 18108351,
					21906769

1813	88090972 (3625, 3626)	1813 88090972 (3625, 3626) Novel Protein sim. GBank	Contains protein domain (PF01417) - glucoamylase	glucoamylase	56182575, 264259, 29331824, 66714117,
		gi 5051536 gb AAD38326.1 AF07372 - (AF073727) EH domain-binding mitalic phosphomotein (Homo coning)	ENTH domain		29331828, 35696052, 264509, 264905,
					254905, 264907, 264908, 66712502, 264909, 1
					203007, 204910, 204391, 204393, 33812038, 3287011, 388011, 384380, 384889, 384784
					203011, 203010, 204700, 204002, 204704, 264603 264360 264366 264368 264360
					2040d3, 2043d9, 2047bb, 2647bB, 2647b9,
					21906/66, 21906/68, 264691, 264693,
					18108374, 35695855, 264634, 264635,
					264637, 264639, 264559, 22279000,
1814		88178047 (3627 3628) Novel Dratein sim CBant ailbeasend (ACOSESSE)			22279002, 264566
		honothetical protein forskidossis thelisasi		UNCLASSIFIED	264488, 35696286, 22278998, 264092,
		יייסיייסיים אייסיים אי			264094, 264259, 29331822, 29331824,
					29331825, 60432289, 29331826, 29331827,
					29331828, 264105, 264107, 52644045,
					56182435, 265009, 60432229, 60433356,
					87168474, 87168559, 264369, 264288,
					21906765, 35695917, 265021, 265022,
					33657023, 33657109, 18108374, 35696423.
					264638, 56526486, 264482
1815	85296473 (3629, 3630)	85296473 (3629, 3630) Novel Protein sim. GBank gil 117788 splP26770 CYA4_RAT		struct	22278999, 264508, 264509, 264907, 264908
	-	ADENYLATE CYCLASE, TYPE IV (ATP			264910 265011 264760 264766 264634
		PYROPHOSPHATE-LYASE) (ADENYLYL CYCLASE)			264636
_					
1816	83738845 (3631, 3632)	83738845 (3631, 3632) Novel Protein sim. GBank		UNCLASSIFIED	18108394, 18108397, 264509, 264907,
	<u></u>	gi 1176623 sp P41846 YO96_CAEEL - HYPOTHETICAL			264908, 264909, 265009, 264591, 265011
		93.9 KD PROTEIN T20B12.6 IN CHROMOSOME III	•		265017, 264687, 264689, 265022, 264691,
					18108362, 18108368, 18108370, 18108374,
_					18108379, 264635, 264557, 264564, 264567
<u>.</u>	66035266 (3633, 3634)	oousozoo (3533, 3534) Novel Protein sim. GBank gij3766377 jemb CAA21429] -	Contains protein domain (PF00400) - histone		264488, 22278997, 22278999, 60432049,
		(AL031907) hypothetical protein (Schizosaccharomyces	WD domain, G-beta repeat		29331822, 29331824, 60432289, 52644045.
		pombe)			60170831, 265017, 265018, 265019,
					18108351, 264682, 52644229, 21906765,
					21906767, 21906768, 52644150, 33657023,
					33657109, 27486262, 18108370, 18108374,
					60170394, 56182323, 22279002
8181	85806775 (3635, 3636)	83806775 (3635, 3636) Novel Protein sim. GBank gij3879121 jemb[CAA94370] -	Contains protein domain (PF00023) - Iranscriptfactor		35696286, 60433356, 264758, 264369,
		(Z70310) predicted using Genefinder; Similarity to Mouse	Ank repeat		264686, 21906769, 264693, 264632
		Composition (FIR Acc. No. 35/1/1); CDNA EST EMBL: 101923			
	- V				
	=•	from this gene; cDNA EST EMBL:D32723 comes from this			
1819	87759572 (7637 3638)	87759572 (1617 1618) Namel Dartein sim CBack			
	(000 '100) = 100	1		UNCLASSIFIED	22278995, 22278997, 22278999, 264259,
		gijoos toosiretjink_uus/71.1jpLHrP - tipoma HMGIC tuston		••	29331826, 264908, 265007, 265008, 265009,
					60432229, 265017, 265018, 265019, 264448,
-					264288, 21906768, 21906769, 265020,
			-	·	18108381, 18108384, 22279000, 22279002,
					264567

1820	87769455 (3639, 3640)				264905, 264907, 264594
1821					264907, 264768, 263978
1822		91221523 (3643, 3644) Novel Protein sim. GBank gil4884130 emb CAB43272.1 -			22278995, 56994075, 22278996, 22278997.
		(AL050101) hypothetical protein [Homo sapiens]			22278998, 264259, 29331824, 29331825,
					29331826, 35696052, 29331828, 264908,
					29331830, 60170831, 264591, 264593.
					60433356, 264596, 265017, 265019,
					18108351, 264763, 264683, 21906765,
					21906767, 21906768, 21906769, 35695917,
					265020, 265021, 33657023, 18108364,
		-			18108370, 35695855, 22279000, 22279002
1823	85522330 (3645, 3646)			UNCLASSIFIED	264488, 264259, 264511, 264288, 264768,
					264693, 35696423, 264634, 18108385,
					264486
1824	_	86612025 (3647, 3648) Novel Protein sim. GBank gil477072 pir 148018 - mucin 7	Contains protein domain (PF00047) - UNCLASSIFIED		264907, 264908, 264909, 264511, 264631,
		precursor, salivary - human	Immunoglobulin domain		264634, 264635, 264637, 264638, 264639,
					264758, 264568
1825		87430125 (3649, 3650) Novel Protein sim. GBank gij3036803 emb CAA18493 - (AL022373) hypothetical protein [Arabidopsis thaliana]		UNCLASSIFIED	60432049, 264910, 264487
1826		91723612 (3651, 3652) Novel Protein sim. GBank		ATPase_associated	ATPase_associated 52644507, 52645156, 52646842, 22278994.
		gil4680685 gb AAD27732.1 AF13295 - (AF132957) CGI-23			22278996, 56994075, 264259, 60432049,
		protein [Homo sapiens]			52645080, 35696052, 66712502, 52644045,
					265008, 265009, 60432229, 60433356,
					60433438, 52646317, 52644296, 265011,
					87168559, 264448, 264288, 264369, 264688,
					52644229, 264689, 21906765, 21906768,
					265020, 60170615, 52644150, 33657023,
					27486262, 27486264, 27486265, 35695763,
					35696423, 35695855, 83373044, 87168518,
					264404, 22279002
1827	1827 81647212 (3653, 3654)				264758

264488, 52646842, 56182575, 22278996, 35696286, 22278997, 22278996, 264093, 60432049, 26278999, 264093, 60432049, 264259, 29331822, 29331824, 66714117, 29331825, 60432289, 29331825, 264302, 26331827, 2644045, 264508, 264507, 66712502, 52644045, 56182435, 265009, 245511, 24512, 265007, 265008, 26509, 60170831, 6043229, 264593, 60433358, 60433438, 264758, 33109954, 21906754, 87168474, 265010, 265011, 87168559, 265017, 265019, 264761, 264762, 264448, 264764, 264683, 264288, 264369, 18108355, 264768, 18108357, 18108358, 264689, 35695749, 265021, 60170615, 33657023, 33657249, 263972, 263072, 2630723, 2655749, 263972, 2630723, 2655749, 263972, 2630723, 2655749, 263972, 2630723, 2655749, 263972, 2630723, 2655749, 263972, 2630768, 264689, 263972, 2630724, 265074, 2630724, 265074, 2630724, 263	25415/6, 35695855, 264635, 264555, 264556, 264558, 264558, 264558, 2645618, 264562, 264564, 26	264908, 265009, 264910 52645156, 22276994, 22278995, 35696286, 2278996, 22278997, 22278998, 22278999, 29331822, 29331825, 35696052, 52646317, 52644296, 81166559, 265019, 21906769, 21906768, 21906767, 21906769, 21906769, 35695917, 265021, 33657023, 52645129, 33657109, 33657182, 27486261, 27486262, 35695763, 263974, 35696423, 35695855,	
oncogene		nuclease	UNCLASSIFIED
Contains protein domain (PF00113) - oncogene			
1828 95074017 (3655, 3656) Novel Protein sim. GBank gil4503571[ref]NP_001419.1 pENO1 - enolase 1, (alpha)		94312942 (3659, 3660) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73, contains large complex repeat CR 73 [Kaposi's sarcomaassociated herpesvirus]	94138063 (3661, 3662) 84521663 (3663, 3664) Novel Protein sim. GBank gil1330345 (U58755) - coded for by C. elegans cDNA yk34b1.5; coded for by C. elegans cDNA yk4668.5; coded for by C. elegans cDNA yk4668.5; coded for by C. elegans cDNA yk4665.5; coded for by C. elegans cDNA yk4665.6; coded for by C. elegans cDNA yk4665.6; coded for by C. elegans cDNA yk43c2.5; coded for by C. elegans cDNA
95074017 (3655, 3656)	80197720 (3657, 3658)		94138063 (3661, 3662) 84521663 (3663, 3664)
1828	1829		1832

<u>6</u>	95314184 (3665, 3666)	1833 95314184 (3665, 3666) Novel Protein sim. GBank gi 5174413 ref NP_006026.1 pCDC4 - CDC42-binding protein kinase beta (DMPK-like)	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	Kinase	22278994, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 60432289, 29331827, 35686052, 2946499, 264508, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 265011, 265017, 264604, 265018, 265019, 264681, 18108351, 264683, 264288, 264685, 21906765, 21906766, 21906766, 21906766, 21906769, 29148629, 265021, 264690, 264692, 33657023, 65274791, 264634, 264635, 264556, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264557, 264559, 284566, 264559, 264559, 264559, 264559, 264559, 264563
06779009	(3007, 3008)				264259, 264907, 264689, 22279000. 22279002
94135718	94135718 (3669, 3670)			UNCLASSIFIED	22278998, 29331822, 29331826, 87168474, 264603, 21906768, 263976, 35695855, 83373044
87348450	(3671, 3672)	.1pUCP4 - uncoupling protein 4	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	ransport	29331825, 264908, 265019, 264764, 264686, 21906765, 264635
94234297 (3673, 3674)	7 (0	Contains protein domain (PF00443) - ubiquitin Ublquitin carboxyl-terminal hydrolase family 2	ubiquilin	22278995, 29146499, 265006, 265008, 265009, 265009, 265010, 264683, 21906765, 29148627, 29148629, 265020, 265021, 265022, 65274620, 18108370, 18108374, 264556, 18108385
94324369 (3675, 3676)	-	Contains protein domain (PF00568) - WH1 domain		2931822, 265017, 264760, 265020, 83373044
87456508	(3677, 3678)	87456508 (3677, 3678) Novel Protein sim. GBank gi[2117310]emb CAB09116.11- (295620) hypothetical protein [Schizosaccharomyces pombe]		UNCLASSIFIED	60433438, 264601, 21906765, 21906766, 265021, 33657109, 264556
87391708	(3679, 3680)	87391708 (3679, 3680) Novel Protein sim. GBank gi]127560 sp P23249 MV10_MOUSE - PROTEIN MOV-10	1	UNCLASSIFIED	264693
85818445	(3681, 3682)	Novel Protein sim. GBank gil4572464[gb]AAD23834.1 AF12365 - (AF123653) FEZ1 [Homo sapiens]			56182575, 29331824, 29331826, 60433356. 264764, 264288, 33657023, 263967, 18108370, 18108374, 264631, 264555, 284556, 264639

65274572, 29331822, 29331824, 29331825, 65714117, 29331826, 29331827, 264907, 264909, 5264909, 5264909, 5265006, 265000, 265000, 265000, 265000, 265000, 265001, 1810836, 264769, 3955702, 264693, 18108364, 284769, 3365702, 264628, 55810764, 56182323, 18108384, 264563, 264564, 264663, 264564, 264663, 264564, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 264664, 26464, 26464, 264664, 264664, 264664, 264664, 26464, 26			264259, 29331824, 264907, 264908, 66712502, 264510, 265007, 265008, 55812038, 265018, 21906765, 52644150, 33657109, 264555, 264556, 264557, 56182323, 18108382, 83373044, 18108385, 264564		22278997, 264259, 29331824, 264909, 18108351, 263974, 22279002	29331825, 29331826, 29331827, 265017, 264683, 264288, 264766, 264768, 21906767, 21906768, 264692, 22279002	iated 56182575, 265018 60432049, 264908	29331824, 35696052, 264910, 60433438, 264688, 35695917, 265020, 52644150, 65274620, 52644332
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transcriptfactor	UNCLASSIFIE	transcriptfactor	tm7	ATPase_assoc struct	dna_ma_bind
			Contains protein domain (PF00628) - transcriptfactor PHD-finger	Contains protein domain (PF00008) - UNCLASSIFIED EGF-like domain	Contains protein domain (PF00010) - transcriptfactor Helix-loop-helix DNA-binding domain		Contains protein domain (PF00008) - ATPase_associated EGF-like domain CPF00169) - Istruct	PH domain Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type
90992645 (3683, 3684) Novel Protein sim. GBank gi 1326268 (U58728) - C54H2. 1 gene product [Caenorhabditis elegans]	5, 3686)	8744764 (3687, 3688) Novel Protein sim. GBank gi 2496887 sp Q09232 YQ22_CAEEL - HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III	95096673 (3689, 3690) Novel Protein sim. GBank gif1175494[sp Q09819 YAC5_SCHPO - HYPOTHETICAL 45.0 KD PROTEIN C16C9.05 IN CHROMOSOME I	84287872 (3691, 3692) Novet Protein sim. GBank gij3881080jembjCAA21739j - (AL032657) similar to EGF-like domain; cDNA EST yk299a12.3 comes from this gene; cDNA EST EMBL:D35398 comes from this gene; cDNA EST yk331h6.5 comes from this gene; cDNA EST yk299a12.5 comes from this gene; cDNA EST yk467q8.5 c	87821497 (3693, 3694) Novel Protein sim. GBank gi 5059323 gb AAD38967.1 AF15152 - (AF151522) hairy and enhancer of split related-1 [Homo sapiens]	86789360 (3695, 3696) Novel Protein sim. GBank gi 5701854 emb CAB52191.1 - (AJ245417) G5b protein [Homo sapiens]	84287874 (3697, 3698) Novel Protein sim. GBank gif4503665[ref]NP_001989.1 pFBLN - fibulin 2 precursor 86689650 (3699, 3700) Novel Protein sim. GBank gif4589582 dbj BAA76813.1 -	(AB023186) KIAA0969 protein [Homo sapiens] 95419789 (3701, 3702) Novel Protein sim. GBank gi[220637 dbj BAA01477] (D10627) zinc finger protein [Mus musculus]
1842 90992645 (3683,	1843 95292692 (3685, 3686)	1844 87444764 (3687,	1845 95096673 (3689	1846 84287872 (3691	1847 87821497 (3693	1848 86789360 (3695	1849 84287874 (3697 1850 86689650 (3699	

n. GBank - 006090.1	=	pPIAS - protein inhibitor of		UNCLASSIFIED	56182575, 35696286, 22278996, 22278997, 22278999, 264490, 60432049, 264259,
activated STAT3	activated STAT3				29331822, 29331824, 29331825, 29331826, 60432289, 29331827, 35696052, 52644045, 265007, 264910, 60432229, 60433356, 60433438, 55812038, 65274444, 265018, 265019, 18108351, 264448, 264686, 264687, 21906767, 21906769, 265021, 265022, 52644150, 264693, 33657109, 18108370, 18108370, 18108370, 264663
91222267 (3705, 3706) Novel Protein sim. GBank gij854065jemb CAA58337j. (X83413) U88 [Human herpesvirus 6]				UNCLASSIFIED	264687, 264768, 52644507, 264769, 21906765, 21906765, 21906778, 21906768, 22278995, 56994075, 22278999, 52644150, 264259, 264692, 29331822, 29331824, 52645129, 29331827, 33656970, 33657349, 35695763,
<u>, </u>	ļ				264508, 264906, 264628, 264907, 264629, 264909, 35696423, 35695855, 264510, 265006, 264511, 264512, 264630, 265009, 264631, 264634, 264636, 264637, 264539, 264638, 264639, 33657402, 18108365, 264602, 265017, 22279000, 265018, 264760, 264762, 264682, 26448, 264764, 264684, 264567, 264288, 264369, 264766, 264567, 264288, 264369, 264468, 264764, 264684, 264567, 264288, 264369, 264566
86038152 (3707, 3708) Novel Protein sim. GBank gi 2072964 (U93569) - putative prise	Novel Protein sim. GBank gi 2072964 (U93569) - putati p150 [Homo sapiens]	, ve		nuclease	264592
91221459 (3709, 3710) Novel Protein sim. GBank gil4539520jembjCAB39994.1 - (AL035424) dA22D12.1 (novel protein similar to Drosophila Kelch (Ring Canal protein, KEL) and a heterogenous set of other types of proteins) [Homo sapiens]	Novel Protein sim. GBank gil4539520jemblCAB39994. (AL035424) dA22D12.1 (novel protein similar to Drosop Kelch (Ring Canal protein, KEL) and a heterogenous se other types of proteins) [Homo sapiens]		Contains protein domain (PF01344) - nucl_recpt Kelch motif		18108392, 52646365, 65274572, 56182575, 22278994, 56994075, 22278996, 35696286, 22278999, 264908, 264510, 265007, 265008, 265009, 264596, 21906754, 87168474, 265011, 87168559, 264681, 264288, 264768, 18108359, 21906764, 21906768, 29148627, 265020, 265021, 265022, 52644150, 33657023, 33657109, 18108372, 18108374, 18108376, 35696423, 264631, 264636, 18108374, 18108381,
94231871 (3711, 3712) Novel Protein sim. GBank gi[3954978 emb CAA06945 - (AJ006278) acetylglucosaminyltransferase-like protein [Musmusculus]	Novel Protein sim. GBank gi[3954978]emb CAA06945 (AJ006278) acetylglucosaminyltransferase-like protein musculus]	Mus		UNCLASSIFIED	264482 56994075, 264259, 29331828, 264511, 264910, 264758, 264693, 264637, 18108381, 83373044
94324455 (3713, 3714) Novel Protein sim. GBank gil4322670[gblAAD16120] - (AF094508) dentin phosphoryn [Homo sapiens]	Novel Protein sim. GBank gil4322670[gb]AAD16120] - (AF094508) dentin phosphoryn [Homo sapiens]			ATPase_associated	ATPase_associated 22278999, 264259, 264906, 60170831, 264448, 264686, 265020, 265022, 33657109. 60170394, 83373044

1858	87628311 (3715, 3716)	1858 87628311 (3715, 3716) Novel Protein sim. GBank gil4981903[gb AAD36415.1 AE00178 - (AE001788) ribosomal protein S15 (Thermotoga maritina)	Contains protein domain (PF00312) - ribosomalprot Ribosomal protein S15	ribosomalprot	264757
1859	84407464 (3717, 3718)	84407464 (3717, 3718) Novel Protein sim. GBank gil4240317[dbjlBAA74937.1] - (AB020721) KIAA0914 protein [Homo sapiens]			22278996, 29331824, 265007, 33109954, 265019, 264369, 21906768, 29148784, 27486261, 52644332, 22279002
1860	17929308 (3719, 3720)	17929308 (3719, 3720) Novel Protein sim. GBank gil4009522 (AF099731) - connexin 31.1 [Homo sapiens]			265019
1861	88086370 (3721, 3722)	88086370 (3721, 3722) Novel Protein sim. GBank gil2143637 pir 184505 - catcium- Contains protein domain (PF00285) dependent actin-binding protein - rat Citrate synthase	Contains protein domain (PF00285) - Citrate synthase		264887, 264259, 29331822, 29331824, 29331825, 265007, 265009, 264591, 33109954, 265010, 265019, 264369, 264288, 264886, 264691, 264693, 27486264, 18108370, 18108374, 263977, 55811576, 56182323, 264639, 22279000, 22279002,
1862	87372923 (3723, 3724)	87372923 (3723, 3724) Novel Protein sim. GBank gij 125493 sp P07313 KMLC_RABIT - MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK)	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	35696286, 264259, 87168474, 264369, 21906766, 264558, 264563
1863		85775037 (3725, 3726) Novel Protein sim. GBank gij3820909 emb CAA09299 - (AJ010842) Dof protein (Drosophila melanogaster)		UNCLASSIFIED	264601, 264766, 29148627, 29148629, 264692, 264629, 264635
1864		85547832 (3727, 3728) Novel Protein sim. GBank gil4322263 gb AAD15985 - (AF077738) metallocarboxypeptidase CPX-1 [Mus musculus]	Contains protein domain (PF00754) - synthase F5/8 type C domain	synthase	22278999, 264259, 264907, 265018, 18108370, 264634, 264635, 264555, 264556, 264638, 18108387
1865		87740827 (3729, 3730) Novel Protein sim. GBank gi 2495727 sp Q93073 Y256_HUMAN - HYPOTHETICAL PROTEIN KIAA0256	·		22278999, 264490, 29331822, 66714117, 66712502, 265006, 265007, 265008, 265009, 264591, 60433438, 265010, 265019, 264760, 264448, 264768, 29148627, 29148629, 265020, 265020, 265027, 18108385, 60432113
1866		87266816 (3731, 3732) Novel Protein sim. GBank gij5262617 emblCAB45748.1 - (AL080157) hypothetical protein [Homo sapiens]		kinase	18108374, 264769, 18108377, 21906765, 21906765, 35696423, 56182575, 21906769, 29148629, 35696423, 56182575, 21906769, 29148629, 35696286, 35695917, 25521, 264510, 264511, 264512, 264534, 264535, 264561, 264557, 23657023, 26331824, 18108385, 21906754, 33657182, 29331824, 18108385, 21906754, 33657182, 265017, 60431802, 22279000, 264682, 264567, 18108372, 264765, 26486
1867		84579159 (3733, 3734) Novel Protein sim. GBank gij3859930 (AF078096) - forkhead/winged helix-like transcription factor 7 [Homo sapiens]		UNCLASSIFIED	264094

264569, 264488, 35696286, 56994075. 264259, 29331822, 29331824, 29331825, 35696052, 29331828, 29146498, 264508, 264509, 264905, 264906, 264907, 264908, 264910, 33657402, 264594, 264758, 55812038, 265011, 264602, 264760, 18108351, 264762, 264681, 264682, 264764, 264369, 21906766, 265681, 264693, 264769, 21906766, 264687, 264768, 264769, 21906766, 264697, 264769, 264629, 18108374, 35696423, 284634, 264635, 264555, 264638, 83373044, 18108385, 56526486, 87168518, 264563, 264564, 264566, 264486, 264567,	264905, 264907, 264908, 265007, 264565, 264566	264766, 264691, 264692, 83373044	264509, 264905, 264908, 264909, 264910, 264762, 264687, 33657023, 264632	264908, 21906766, 18108370, 263974, 87168518	264908, 264910, 87168559, 21906766. 264636
	UNCLASSIFIED 26	UNCLASSIFIED 26	UNCLASSIFIED 26	26	
Contains protein domain (PF01428) - ubiquitin AN1-like Zinc finger					Contains protein domain (PF00412) · homeobox LIM domain containing proteins
1878 95351056 (3755, 3756) Novel Protein sim. GBank gil4510345[gb]AAD21434.1 - (AC006921) unknown protein [Arabidopsis thaliana]	95310883 (3757, 3758) Novel Protein sim. GBank gil4929643 gblAAD34082.1 AF15184 - (AF151845) CGI-87 protein [Homo sapiens]	91012978 (3759, 3760) Novel Protein sim. GBank gil1550785 emb CAA69283 - (708026) Immune associated protein 38 (Mus musculus)	80214949 (3761, 3762) Novel Protein sim. GBank gij93144 pir B40505 - hypothetical protein - suid herpesvirus 1 (strain Indiana- Funkhuser or Becker)	86582450 (3763, 3764) Novel Protein sim. GBank gil2384956 (AF022985) - No definition line found (Caenomabditis elegans)	IN 2 (TES2)
95351056 (3755, 3756)	95310883 (3757, 3758)	91012978 (3759, 3760)	80214949 (3761, 3762)	86582450 (3763, 3764)	94216817 (3765, 3766)
1878	1879	1880	1881	1882	1883

264488, 18108394, 56181686, 22278998, 60432049, 264259, 29331822, 29331824, 29331824, 29331825, 66714117, 29331827, 29331824, 29331825, 66714117, 29331827, 256906, 264508, 264509, 264906, 264907, 264509, 264906, 264910, 264511, 264908, 264907, 264501, 264511, 264600, 265010, 264510, 264501, 264600, 265010, 264010, 265019, 264605, 264600, 265017, 264604, 265019, 264605, 264681, 264769, 264761, 264682, 264769, 29148784, 264929, 264690, 264911, 33657023, 264692, 264691, 33657023, 264692, 264631, 33657023, 264692, 264634, 264634, 264634, 65810764, 55811576, 35696423, 35598855, 264638, 56182332, 264639, 18108372, 264638, 56182332, 264639, 18108382, 83373044, 18108385, 60432113, 22279002, 264567, 264565, 264566, 264486, 264567, 264567, 264567, 264567, 264567, 264567, 264565, 264566, 264486, 264567, 264565, 264566, 264486, 264567, 264567, 264565, 264566, 264486, 264567, 264565, 264566, 264486, 264567, 264565, 264566, 264486, 264567, 264567, 264567, 264486, 264567, 264565, 264566, 264486, 264567, 264567, 264567, 264486, 264567, 26	56182575, 264259, 264905, 264909, 265008, 264596, 264766, 265020, 264628, 60431528, 264634, 56526486, 264080, 26456, 264080, 26458	22278998, 22278999, 60432049, 264910, 265018, 264766, 21906768, 29148629, 264628, 264586, 264628, 264688, 264628, 264628, 264628, 264628, 264628, 264628, 264628, 264688, 264628, 264628, 264628, 264628, 264628, 264628, 264628, 264688, 264628, 264628, 264628, 264628, 264628, 264628, 264628, 264688, 264628, 264688, 264628, 264688, 2646	29311822, 29331824, 29331825, 29331820, 29331827, 264906, 265007, 264681, 264768, 29148627, 264693, 18108364, 35696423, 65274791, 35695855, 264632, 56182323, 264639, 264563	263978	264906, 55812038, 264758, 265010, 265018, 265019, 18108351, 264288, 21906765, 21906769, 265021, 33657023, 33657109, 56182323, 83373044, 18108385, 22279000, 22279002	18108398, 29147620, 264907, 265009, 264600, 265018, 18108351, 264288, 264689, 21906765, 21906768, 21906769, 264691, 264692, 264693, 264628, 18108370, 264636, 264558, 264504
	UNCLASSIFIED		ATPase_associated		helicase	UNCLASSIFIED
			Contains protein domain (PF00168) - ATPase_associated	Contains protein domain (PF01454) - MAGE family	Contains protein domain (PF00176) - helicase SNF2 and others N-terminal domain	
Novel Protein sim. GBank gil4829643 gb AAD34082.1 AF15184 - (AF151845) CGI-87 protein [Homo sapiens]	87644280 (3769, 3770) Novel Protein sim. GBank gil2507155 sp P37370 VRP1_YEAST - VERPROLIN	86674062 (3771, 3772) Novel Protein sim. GBank gil2854158lgblAAC02577.1 - (AF045641) No definition line found [Caenorhabditis elegans]	94139139 (3773, 3774) Novel Protein sim. GBank gi 5174421 ref NP_006023.1 pCPNE - copine VI (neuronal) C2 domain	87822804 (3775, 3776) Novel Protein sim. GBank gij3319931 emb CAB10841 - (298046) dJ14O9.2 (Melanoma-Associated Antigen MAGE LIKE) [Homo sapiens]	91255783 (3777, 3778) Novel Protein sim. GBank gij 1083308jpir A56559 - enhancer-trap-locus-1 protein - mouse (fragment)	87626705 (3779, 3780) Novel Protein sim. GBank gil4240195 db BAA74876.1 - (AB020660) KIAA0853 protein [Homo sapiens]
1884 95310885 (3767, 3768) Novel Protein sim. GBank gip422643)gbpAnD34082. protein [Homo sapiens]	1885 87644280 (3769, 3770)	1886 86674062 (3771, 3772)	1887 94139139 (3773, 3774)	1888 87822804 (3775, 3776)	1669 91255783 (3777, 3778)	1890 87626705 (3779, 3780)

1691				UNCLASSIFIED	264686, 264768, 264687, 264692, 264693, 29331822, 29331824, 264508, 264905, 264906, 18108370, 264508, 264907, 264908, 264909, 18108379, 255007, 265008, 264910, 264632, 264591, 264601, 264605, 264563, 265010, 265011, 264601, 264605, 264563, 264369
1892	87642825 (3783, 3784)	Novel Protein sim. GBank gil5689535 dbj BAA83051.1 - (AB029022) KIAA1099 protein [Homo sapiens]	Contains protein domain (PF01412) - Putative GTP-ase activating protein for Arf	struct	22278995, 264509, 87168559, 18108351, 264448, 264682, 265020, 264693, 18108374, 2279000
1893	88533826 (3785, 3786)			laminin	264569, 65274572, 22278997, 22278999, 264259, 29331824, 66714117, 29331826, 265006, 265008, 265009, 264592, 265018, 264681, 26448, 264683, 18108354, 264369, 284684, 264685, 264766.
					264687, 264689, 21906768, 255020, 255022, 60170615, 52644150, 264690, 264681, 264692, 33657023, 264690, 33657109, 264632, 18108374, 3569855, 264630, 264634, 264537, 264558, 60170394, 18108381, 18108388, 22279000
1894					264508, 264905, 264906, 264997, 264594, 264684, 264690, 264692, 264630, 264635, 264636, 264639, 264563
1895		87631891 (3789, 3790) Novel Protein sim. GBank gi 5262574 emb CAB45729.1 - (AL080133) hypothetical protein [Homo sapiens]	Contains protein domain (PF00435) - Spectrin repeat		56182575, 264259, 60432289, 29331826, 264107, 264905, 264908, 264910, 60170831, 264758, 265010, 265018, 264448, 264288, 264768, 33657109, 264628, 55810764, 18108379, 264634, 56182323, 56556486
1896			Contains protein domain (PF00627) - UNCLASSIFIED UBA domain	UNCLASSIFIED	264907, 265008, 264682, 264686, 21906768, 264629, 264631, 264634, 264555
1897		80565569 (3793, 3794) Novel Protein sim. GBank gij728836 sp P39193 ALU6_HUMAN - !!!! ALU SUBFAMILY SP WARNING ENTRY !!!!		cadherin	264259
1898	87617637 (3795, 3786)	87617637 (3795, 3796) Novel Protein sim. GBank gi1127560 sp P23249 MV10_MOUSE - PROTEIN MOV-10		helicase	22276996, 22278998, 22278999, 29331824, 29331825, 60432289, 29331827, 35696052, 29331828, 265008, 265019, 264681, 264682, 264448, 264389, 52644229, 21906765, 21906768, 21906769, 60170615, 55810764, 22279000
1899		86673097 (3797, 3798) Novel Protein sim. GBank gi 2909819 (AF031548) - erythrocyte membrane glycoprotein Rh50 [Homo sapiens]	Contains protein domain (PF00909) - glycoprotein Ammonium Transporter Family	glycoprotein	264259, 264508, 264909, 60432229, 264769, 21906765, 21906769
1900		87641858 (3799, 3800) Novel Protein sim. GBank gil4102881 (AF017250) - vitellogenin precursor (Oreochromis aureus)		UNCLASSIFIED	264683

264488, 52644507, 52645156, 18108396, 52646365, 52646842, 18108397, 56182575, 22278994, 22278995, 56994075, 35696286, 22278997, 22278995, 56994075, 35696286, 22278997, 22278998, 264490, 60432049, 264259, 29331827, 29331826, 264300, 29331827, 29331828, 3369052, 33656970, 29416498, 264907, 29331830, 264908, 5264045, 264907, 29331830, 264908, 5264045, 264907, 29331830, 264908, 5264045, 264907, 29331830, 264908, 264709, 264907, 29331830, 264908, 52644045, 264907, 29306754, 3365702, 264907, 265019, 265011, 87168559, 265017, 265018, 265019, 18108351, 264428, 265917, 265018, 265021, 265021, 52644150, 18108362, 33657023, 264632, 263917, 265020, 265021, 52644150, 18108362, 33657023, 264632, 263981, 52644332, 60431850, 264638, 263981, 52644332, 60431850, 264634, 18108358, 87168518, 60432113, 264564		D 264637
transport	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00242) - Iransport DNA polymerase (viral) N-terminal domain	Contains protein domain (PF00293) - Bacterial mufT protein	
1901 95196647 (3801, 3802) Novel Protein sim. GBank gil585959jsp P38378j561A_RAT PROTEIN SEC61 ALPHA SUBUNIT	(AF098796) SLM-1 [Mus musculus] (AF098796) SLM-1 [Mus musculus] (AF098741) - unknown [Rattus norvegicus] 80434213 (3807, 3808) Novel Protein sim. GBank gij3747107 (AF095741) - gij1352911[sp[P47147]YJ80_YEAST - HYPOTHETICAL 80.2 KD PROTEIN IN CPA2-NNF1 INTERGENIC REGION 95351140 (3809, 3810) Novel Protein sim. GBank gij3043714[dbj[BAA25521] - (AB011167) KIAA0595 protein [Homo sapiens]	
95196647 (3801, 3802)	87778554 (3805, 3806) 80434213 (3807, 3808) 95351140 (3809, 3810)	12763822 (3811, 3812)
1901	1903	8

1911	91725345 (3821, 3822)	91725345 (3821, 3822) Novel Profein sim, GBank	Contains protein domain (PF01119) -Inuclease	nuclease	18108394, 56182575, 56182181, 29331826.
		gil4809339lgblAAD30184.1 AC00653 - (AC006530)	DNA mismatch repair protein		29331827, 33656970, 264906, 265007.
		hypothetical protein [Homo sapiens]			264591, 55812038, 87168559, 264448,
					264369, 21906765, 21906768, 265022,
				_	264691, 264693, 18108365, 55811576,
					264556, 18108385, 18108388
1912		95413519 (3823, 3824) Novel Protein sim. GBank gil5689439 dbj BAA83003.1 -	Contains protein domain (PF00098) - UNCLASSIFIED	UNCLASSIFIED	18108397, 56182575, 56181686, 22278994,
		(AB028974) KIAA1051 protein [Homo sapiens]	Zinc finger, CCHC class		22278995, 56994075, 22278996, 22278999,
					264259, 29331822, 29331824, 56182181,
					29331825, 66714117, 35696052, 264905.
					264906, 264907, 264908, 52644045.
					56182435, 265007, 265008, 264910, 265009,
					264591, 264596, 65274444, 55811386,
					87168474, 265011, 87168559, 265018,
					265019, 264760, 18108351, 264681, 264369,
					264684, 264288, 264686, 264768, 21906765,
					21906766, 21906767, 21908768, 21906769,
					55811957, 265020, 265021, 265022,
					60170615, 264692, 33657023, 264693,
					18108376, 55811576, 35696423, 65274791,
					264637, 56182323, 83373044, 56526486,
					22279002, 264563, 264566
1913	95305546 (3825, 3826)	95305546 (3825, 3826) Novel Protein sim. GBank			56182575, 22278994, 22278995, 56994075,
	•	gi 5032245 ref NP_005665.1 pZNF2 - zinc finger protein			22278996, 22278998, 22278999, 29331826,
		(C2H2) homologous to mouse MOK-2			29331827, 265006, 55812038, 265010,
			٠		265017, 265018, 265019, 264681, 18108351,
					264683, 264764, 264369, 264288, 264685,
					264686, 264769, 21906765, 21906766.
	•				21906768, 21906769, 55811957, 265020,
					265022, 264691, 55811576, 264634, 264635,
					264638, 56182323, 83373044, 18108385
1914	83423982 (3827, 3828)	83423982 (3827, 3828) Novel Protein sim. GBank gil4589604 dbj BAA76824.1 -	Contains protein domain (PF00036) - struct	struct	56182575, 29331824, 35696052, 264906,
		(AB023197) KIAA0980 protein [Homo sapiens]	EF hand		264908, 264828, 264909, 264592, 264758,
					87168559, 18108351, 18108354, 264684.
					264686, 33657023, 264693, 264628, 264631,
					264632, 264634, 264635, 264639
1915		95340459 (3829, 3830) Novel Protein sim. GBank gij5689415[dbj]BAA82991.1 -		UNCLASSIFIED	264259, 29331824, 29331826, 29331827.
_		(AB028962) KIAA1039 protein [Homo sapiens]			264508, 264909, 265009, 265017, 265019,
					264768, 264769, 264689, 264628, 264635,
					264637, 264639, 83373044, 264565
1916	1916 79640761 (3831, 3832)				264693, 264639

264769, 264689, 21906765, 21906768, 22278996, 264259, 264691, 264693, 29331824, 29331825, 29331826, 29331828, 264905, 264906, 264628, 264907, 264908, 264909, 264630, 264910, 264634, 264635, 264638, 263981, 264639, 264758, 18108385, 21906754, 265011, 264604, 264563, 18108351, 264762, 264763, 264764, 264766	264488, 18108392, 18108357, 21906765, 21906767, 21906768, 56182575, 21906769, 22278994, 35696286, 35695917, 22278996, 22278994, 35696286, 3569517, 22278996, 22278997, 265021, 265022, 264534, 264690, 264691, 264692, 33657023, 264693, 264691, 264692, 33657023, 264693, 2645124, 29331825, 2345708, 264906, 264506, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264907, 18108370, 264510, 264909, 18108372, 265008, 264631, 265007, 264934, 264535, 264594, 60170394, 264595, 264594, 60170394, 264595, 264594, 60170394, 264595, 264604, 265010, 265011, 264063, 264604, 265010, 264017, 264603, 264604, 265011, 264605, 264487, 264685, 264488, 264487, 264685, 264487, 264686, 264487, 264686, 264487, 264686, 264487, 264686, 264487, 264686, 264487, 264687, 264687, 264487, 26468	.e 22278997, 29331822, 265007, 60170831, 60432229, 60433438, 264448, 264682, 264288, 55811957, 33657023, 33657109, 65274791, 56182323, 22279002	UNCLASSIFIED 264510, 264511, 264512, 264566	UNCLASSIFIED 264693
Sfrud	Struct	oxidase		
Novel Protein sim. GBank gil5689391 db BAA82979.1 - (AB028950) KIAA1027 protein [Homo sapiens]	95302795 (3835, 3836) Novel Protein sim. GBank gil5281517[gb]AAD41524.1 AF15483 - (AF154831) PV-1 [Raftus norvegicus]	94143847 (3837, 3838) Novel Protein sim. GBank gij3878584 emb CAB01237 - (Z77667) cDNA EST EMBL:C08125 comes from this gene; cDNA EST EMBL:C09753 comes from this gene [Caenorhabditis elegans]	91229953 (3839, 3840) Novel Protein sim. GBank gij1809231 (AC000115) - coded for by human cDNAs R76043 (NID: 9850725), R65857 (NID: 9838495) and H12868 (NID: 9877688) [Homo sapiens]	Novel Protein sim. GBank gil4580997 gb AAD24571.1 AF12108 - (AF121081) CAMP
1917 87821680 (3833, 3834) Novel Protein sim. GBank gil (AB028950) KIAA1027 prote	1918 95302795 (3835, 3836) N	1919 94143847 (3837, 3838) I	1920 91229953 (3839, 3840)	1921 79555226 (3841, 3842) Novel Protein sim. GBank gil4580997[gb AAD24571.1]AF121

oi.	oikidi					mi
264686, 264688, 264490, 18108370, 264909, 18108374, 265008, 264557, 264564, 18108351	264488, 264489, 22278995, 264094, 264258, 35696052, 264509, 264906, 264906, 264906, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264908, 264511, 264512, 265007, 264910, 265009, 264758, 264596, 264769, 264767, 26448, 264769, 264767, 26448, 264764, 264767, 26448, 264768, 264767, 264689, 264689, 264769, 33657023, 264693, 264691, 33657023, 264693, 264634, 264634, 264634, 264634, 264634, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264658, 264567, 264567, 264563, 264565, 264565, 264567, 264567, 264563, 264565, 264566, 264567, 264687, 2	29331826, 264906, 264908, 264595, 265017, 265018, 265019, 265021, 264691, 264693, 264637, 18108385, 264565	22278995, 22278996, 22278997, 264259, 29331824, 66714117, 29146499, 264909, 52644045, 265008, 265009, 264758, 265011, 265017, 264605, 264448, 264288, 264692, 33657109, 18108374, 60170394	264905, 264906, 264907, 264910, 264593, 265018, 264760, 264764, 264288, 264692, 264693, 264693, 264631, 264634, 264637, 264689, 264683	264489, 264259, 265017, 265021, 264692	264508, 264591, 33657402, 265017, 264768, 264632, 264556, 264639
UNCLASSIFIED	UNCLASSIFIED		ribosomalprot	struct	tnf	UNCLASSIFIED
		Contains protein domain (PF00097) - Zinc finger, C3HC4 type (RING finger)	Contains protein domain (PF00312) - ribosomalprot Ribosomal protein S15		Contains protein domain (PF00193) - Inf	
1922 87641863 (3843, 3844) Novel Protein sim. GBank gil138595 sp P02845 VIT2_CHICK - VITELLOGENIN II PRECURSOR (MAJOR VITELLOGENIN) [CONTAINS: LIPOVITELLIN I (LVI); PHOSVITIN (PV); LIPOVITELLIN II (LVII); YGP40]	94323589 (3845, 3846) Novel Protein sim. GBank gi 119110 sp P03211 EBN1_EBV - EBNA-1 NUCLEAR PROTEIN	87338925 (3847, 3848) Novel Protein sim. GBank gil3877655 emb CAA96657 - (Z72511) possible zinc finger protein; cDNA EST EMBL:M89115 comes from this gene; cDNA EST EMBL:D71533 comes from this gene; cDNA EST EMBL:D72314 comes from this gene; cDNA EST EMBL:D75164 comes from this gene; cDNA EST EMBL:C7.1.	87628338 (3849, 3850) Novel Protein sim. GBank gi 4981903 gb AAD36415.1 AE00178 - (AE001788) ribosomal protein S15 [Thermotoga maritima]	88094739 (3851, 3852) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73, Contains large complex repeat CR 73 (Kaposi's sarcoma-associated herpesvirus) 85654857 (3853, 3854) Novel Protein sim. GBank gil3043632 dbjjBAA25480 -	87799054 (3855, 3856) Novel Protein sin. GBank gil665761[dbj BAA13377] - [Opposed Protein sin. GBank gil665761[dbj BAA13377] - [Opposed Bank gil665761[dbj BAA13377] - [Opposed Bank gil665761]	86997236 (3857, 3858) Novel Protein sim. GBank gil5001993[gblAAD37247.1]AF13432 - (AF134321) chimeric AFGPArypsinogen-like serine protease precursor [Dissostichus mawsoni]
87641863 (3843, 3844)	94323589 (3845, 3846)	87338925 (3847, 3848)	, , , , , , , , , , , , , , , , , , ,	88094739 (3851, 3852) 85654857 (3853, 3854)	87799054 (3855, 3856)	86997236 (3857, 3858)
1922		1924		1926	1928	1929

35696286, 29331828, 264905, 264907, 264908, 264908, 264909, 264511, 264910, 264758, 264601, 265017, 265019, 264605, 264760, 264764, 264766, 264688, 264769, 265022, 35698423, 264638, 60432113	264906, 264908, 60432229, 264758, 264764, 264288, 265020, 264692, 264634, 264637	264684, 264691, 264635	264595	56182575, 56182435, 264510, 264757, 264758, 55812038, 55811386, 265018, 55811150, 21908765, 264691, 264631, 264633, 26	264686, 265011, 264511, 264905, 18108351, 264564, 264681, 264259, 18108370, 264566, 264764, 264369, 264595	60432289, 265007, 265010, 265011, 265019, 33657109, 18108374	85658542, 21906767, 35695917, 60170615, 264693, 33657109	264488, 29146498, 264905, 264559	65274572, 2227899, 60432289, 56182435, 60433356, 265017, 21906765, 21906766, 21906768, 55811957, 27486264, 35696423, 60432113, 264564	264488, 29331822, 264448, 264683, 264288, 265020, 33657023, 264631	56994075, 2227898, 264259, 29331824, 29331827, 264905, 265008, 33857084, 265017, 265017, 265018, 264289, 264687, 21906765, 21906766, 21906767, 265020, 52644150, 27486264, 83373044, 18108387, 60432113, 22279002, 264565	264488, 18108398, 29331825, 27486261, 264509, 18108370, 18108374, 264482	22278998, 29331822, 29331827, 35696052, 264511, 265009, 264592, 60432229, 265017, 265018, 265019, 264684, 264692, 33657109, 65274791, 264636
phosphatase	kinase		UNCLASSIFIED	struct	UNCLASSIFIED		kinasa	collagen	UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED	. kinase
	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain						Contains protein domain (PF00023) - kinase Ank repeat	Contains protein domain (PF00888) - collagen Cullin family		Contains protein domain (PF00560) - struct Leucine Rich Repeat			Contains protein domain (PF00400) - kinase WD domain, G-beta repeat
1930 87889128 (3859, 3860) Novei Protein sim. GBank gi 1709230 sp P52963 NBL4_MOUSE - NBL4 PROTEIN	87797279 (3861, 3662) Novel Protein sim. GBank gil404634 (U01840) - serine/threonine kinase IMus musculus)			84426360 (3867, 3868) Novet Protein sim. GBank gil4115748 dbj BAA36494 - (AB022023) nonmuscle myosin heavy chain B [Bos taurus]	((95414338 (3871, 3872) Novel Protein sim. GBank gil4827040 ref NP_005110.1 pTRAP - thyroid hormone receptor-associated protein, 150 kDa subunit		87403277 (3875, 3876) Novel Protein sim. GBank gil4544431lgblAAD22340.1lAC00695 - (AC006955) hypothetical protein [Arabidopsis thaliana]	91004978 (3877, 3878) Novel Protein sim. GBank gil500858 dbj BAA03210 - (D14168) 50kDa lectin [Bombyx mori]	87348810 (3879, 3880) Novel Protein sim. GBank gil1946300 emb CAA73132 - (Y12529) hypothetical protein [Silene latifolia]	94147177 (3881, 3882) Novel Protein sim. GBank gij4206386 (AF060570) - rig-1 protein [Mus musculus]	87641870 (3883, 3884) Novel Protein sim. GBank gil4927204lgb AAD33049.1 AF13391 - (AF133911) ARL-6 interacting protein-4 [Mus musculus]	94325298 (3885, 3886) Novel Protein sim. GBank gij3122952 sp O15736 TIPD_DICDI - TIPD PROTEIN
87889128 (3859, 3860)	87797279 (3861, 3862)	15030972 (3863, 3864)			87752511 (3869, 3870)		_			87348810 (3879, 3880)			
1930	1831	1932	1933	1934	1935	1936	1937	1938	1939	1940	1941	1942	1943

1944	94232958 (3887, 3888)	1944 94232958 (3887, 3888) Novel Protein sim. GBank gil1799570 dbj BAA13432 -		UNCLASSIFIED	65274572, 22278994, 22278995, 22278996,	_
					22278998, 22278999, 264259, 52645080, 29331824, 29331826, 29331827, 66712602	
					56182435, 264512, 264910, 60170831,	
					60432229, 60433356, 33657402, 60433438,	_
					264596, 33109954, 21906754, 87168474,	
					87168559, 265017, 265018, 265019,	
					18108351, 264369, 264686, 264768,	
					21906765, 21906766, 21906767, 21906769,	
					35695917, 265020, 265021, 60170615,	
					264692, 33657023, 18108370, 18108374,	
				4,	35696423, 35695855, 264634, 60170394,	_
					264639, 83373044, 18108385, 56526486,	
1045				ij	87168518, 60432113, 22279000, 264563	
2		ovoviora (Josep, Josep) Novel Protein Sim. GBank		UNCLASSIFIED	264488, 22278996, 264510, 264511,	_
		griesz/zoelgapAdusso49.1 AF13391 - (AF133911) ARL-6 interacting protein-4 [Mus musculus]			18108351, 264683, 264486, 264567	
1946		87443990 (3891, 3892) Novel Protein sim. GBank		UNCLASSIFIED	60432289 29331827 35696052 265007	-
		gi[2498104 sp Q27969 AD50_BOVIN - ADRENAL			265008, 60433356, 60433438, 264369.	_
		MEDULLA 50 KD PROTEIN			56181562, 21906767, 52644150, 264693,	
7,70	17000 00007 00000000				27486264, 264637, 87168518, 264563	
	00430002 (3093, 3094)	obasodoz (soss, sosa) Novel Protein sim. GBank		rnapolymerase	22278998, 264905, 264906, 264908, 264909,	_
		Bilas 1400 (Spinos of the Community of t			264512, 264758, 264762, 264682, 264683,	_
		POLYMERASE I 135 KU POLYPEPTIDE (RNA			264764, 264288, 264768, 264688, 21906768,	
		POLIMERASE 1 SUBUNIT 2) (RPA133) (RNA			264693, 18108374, 35695855, 264635,	
1048	0510017/ /200E 200C)	PULTMERASE 112/ KD SUBUNIT)			264637, 264638, 18108385, 22279002	
	93139114 (3093, 3090)	63.139.174 (3083), 3080) [NOVel Protein sim. GBank gij5420387/emb CAB46679.1]		struct	264909, 60170831, 264591, 264594, 235010.	_
		(AJZ43439) proteopnosphoglycan [Letshmania major]			265011, 264764, 264369, 264689, 264631,	
	7640129 (3897, 3898)				264638	
1950	87788531 (3899 3900)	87788531 (3899-3900) Novel Protein sim CBask April 2008 (1888)			264369	_
_	(2000)	(269637) predicted using Genefinder: Similarity to E coti	Contains protein domain (PF00857) - UNCLASSIFIED Isochorismalase (amily		264488, 264768, 264688, 264689, 264259,	
		hypothetical protein YCAC (SW:YCAC ECOL!)			25331022,3303/109,29331020,204103, 364500 18108370 35605856 364510	
		[Caenorhabditis elegans]			265008, 265009, 33657402, 18108385	
					265018, 264563, 55811150, 18108351	
_					264369, 264288, 18108354	
6	60988253 (3901, 3902)	60988253 (3901, 3902) Novel Protein sim. GBank gi[2626753 db] BAA23424 -	Contains protein domain (PF00916) - transport		56182575, 22278997, 52645080, 29331824,	_
		(Abuso 62) surate transponer [Arabidopsis thaliana]	Sulfate transporter family		29331825, 29331827, 55812038, 52646317,	
					265018, 265019, 264369, 21906765,	
					21906767, 55811957, 265020, 265021,	_
					33657023, 264693, 35695763, 56182323, 22270002	
1952	87069775 (3903, 3904)	87069775 (3903, 3904) Novel Protein sim. GBank	Contains protein domain (PF00106) - reductase		264591 264593 264594 264595 26455	_
		gil4929633lgbJAAD34077.1JAF15184 - (AF151840) CGI-82	short chain dehydrogenase		204356, 264557, 264558, 264565, 264556, 264557, 264558, 264565	
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nt 264259, 264558	264488, 263994, 18108394, 35696286, 22278998, 29331822, 66714117, 29331826, 29331827, 264509, 264905, 264906, 264907, 264908, 264509, 264905, 264906, 264907, 265008, 264910, 265009, 60170831, 6043229, 265011, 265017, 264603, 264604, 265019, 18108351, 264685, 264769, 265019, 18108351, 264685, 264769, 35695917, 265020, 265021, 264691, 33657023, 264692, 33657109, 264628, 18108374, 35696423, 55811576, 35695855, 264530, 264638, 264557, 264639, 264638, 264557, 264569, 264638, 264557, 264569, 264658, 264567, 264639, 264557, 264569, 264658, 264569, 264658, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 264659, 264569, 26469, 264669, 264669, 264669, 264669, 264669, 264669, 264669, 264669, 264669, 26469,		264488, 56182575, 22278936, 22278939, 264259, 29331824, 29331825, 29331824, 29331825, 29331824, 29331825, 29331827, 35696052, 29331824, 29331825, 264906, 264909, 264511, 265007, 264910, 264591, 55812038, 265010, 265018, 18108351, 264768, 56181562, 264689, 21906768, 21906769, 265022, 264691, 264628, 18108374, 55810764, 55811576, 35695855, 264631, 264632, 264635, 264631, 264632, 363373044, 18108385, 22279000, 22279002, 264563.
сотріете	ubiquitin	UNCLASSIFIED	cadherin
Contains protein domain (PF00386) - complement C1q domain	Contains protein domain (PF00443) - ubiquitin Ubiquitin carboxyl-terminal hydrolase family 2		Contains protein domain (PF00028) - cadherin Cadherin domain
1953 20470371 (3905, 3906) Novel Protein sim. GBank gi 1168715 sp P31721 C1QB_RAT - COMPLEMENT C1Q SUBCOMPONENT. B CHAIN PRECURSOR	Novel Protein sim. GBank gi 4240271 db BAA74914.1 - (AB020698) KIAA0891 protein [Homo sapiens]	1	95092121 (3911, 3912) Novel Protein sim. GBank gil1665821 dbj BAA13407 - (D87469) Similar to D.melanogaster cadherin-related tumor suppressor [Homo sapiens]
)470371 (3905, 3906)	3908) (3907, 3908)	1955 95308310 (3909, 3910)	092121 (3911, 3912)
1953 20	4 6 6 6		1856 950 950

1982.1 - UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED Struct		94326510 (3913, 3914)	1957 94326510 (3913, 3914) Novel Protein sim. GBank gil4569674 dbj BAA76856.1 - (AB023229) KIAA1012 protein iHomo sapiensi		UNCLASSIFIED	52646842, 56182575, 22278997, 22278998,	
						227.0999, 29331824, 66714117, 29331827, 29146498, 264593, 33657402, 33109954, 87168474, 265018, 264448, 264369, 264288	
						264766, 21906766, 21908767, 21906768, 21906769	
	_					65274620, 27486264, 33657349, 27486265.	
	_	95313902 (3915, 3916)	Novel Protein cim CBank nild24022714hii04 474802 41			35695855, 22279002, 264482	
	_	(0.00)	[A8020676] KIAA0869 protein [Homo capiene]			22278999, 264092, 264094, 264259,	
	_					60432049, 29331824, 56182181, 66714117,	
	_					264107, 264109, 264909, 264511, 60170831,	_
	-					60432229, 21906754, 265010, 21906769,	
	_					35695917, 265022, 65274620, 263967,	
						263976, 35696423, 264631, 264632, 264634, 264626, 4840836, 20072000, 20072000	
Schizosaccharomyces 1274851 dbj BAA21515		85701470 (3917, 3918)	Novel Protein sim. GBank gi[2281983 emb CAB10860			264593, 265019	
			pombe]				
[Homo sapiens] 13721653 db BAA33581 - eph etase 5 [Rattus norvegicus] eph 15262638 emb CAB45757.1 - cadherin 15262638 emb CAB45757.1 - cadherin 14107017 db BAA38294 - Contains protein domain (PF01428) - ubiquitin 24107017 db BAA38294 - AM1-like Zinc finger		80308608 (3919, 3920)	Novel Protein sim. GBank gi[2274851 dbj BAA21515 -			264905 264906 264907 264909 264909	
UNCLASSIFIED UNCLASSIFIED Eph Eph			(D64159) 3-7 gene product (Homo sapiens)	•		265006, 265007, 264910, 264595, 265017.	
UNCLASSIFIED UNCLASSIFIED Eph Eph						264604, 265018, 18108351, 264764, 264369,	
13721653 db BAA33581 - eph eph						264766, 264768, 21906765, 18108368,	
UNCLASSIFIED UNCLASSIFIED Eph Eph						264629, 18108379, 264635, 264636, 264637,	
UNCLASSIFIED UNCLASSIFIED EPh Eph		16292607 (3921, 3922)				264638, 264486	
3721653 db BAA33581 - etase 5 [Rattus norvegicus] etase 5 [Rattus norvegicus] 5262638 emb CAB45757.1 - 15262638 emb CAB45757.1 - 1610 [Homo sapiens] 4107017 db BAA36294 - AN1-like Zinc finger		91008385 (3923, 3924)			Ī	(04035	_
efase 5 [Rattus norvegicus] efase 5 [Rattus norvegicus] [5262638[emb[CAB45757.1] - cadherin tein [Homo sapiens] [4107017[dbj BAA36294] - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger		90936017 (3925, 3926)	Novel Protein cim CBank all 2221652145:104 4225841		ASSIFIED	55274572, 264592, 264593, 265019, 264691	
		(0.50, '0.50)	[Novel Flores] Sint. Obank gils/2.1833[db] BAA33381] - [(AB012933) acvi-CoA synthetase 5 [Rathis populations]			5274572, 18108398, 35696286, 29331825,	
5262638 emb CAB45757.1 - cadherin tein [Homo sapiens]	_ `		ליים ביים ליים ביים ליים ביים ביים ביים			10432289, 29331827, 264828, 265006,	_
5262638 emb CAB45757.1 - cadherin cadherin	-					65009, 60433356, 60433438, 21906754,	_
5262638 emb CAB45757.1 - cadherin lein [Homo sapiens]						(65020, 265021, 33657023, 33657109, 1248626, 266066, 26466	_
Itein [Homo sapiens] 107017[dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger	6	4317605 (3927, 3928)	Novel Protein sim. GBank gi 5262638 emb CAB45757.1 -			64488 264092 264259 264509	
 4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi AN1-like Zinc finger			(AL080169) hypothetical protein [Homo sapiens]			154906 264907 264908 264909 264640	
 4107017 dbj BAA36294 - Contains protein domain (PF01428) - Ubiquitin avignyi AN1-like Zinc finger						64511, 265007, 265009, 264910, 264592	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger						64593, 264594, 264595, 264758, 264600	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi AN1-like Zinc finger		-				64603, 264604, 264605, 264760, 264762,	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger						264448, 264764, 264288, 264685, 264766,	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger						264768, 264769, 21906766, 264691, 264692,	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi] AN1-like Zinc finger						64693, 18108370, 264628, 264629,	
						8108374, 264630, 264631, 264634, 264636,	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - Ubiquitin avignyi] AN1-like Zinc finger						64637, 264638, 18108382, 83373044,	
4107017 dbj BAA36294 - Contains protein domain (PF01428) - ubiquitin avignyi					<u>- 0</u>	8108385, 264483, 264564, 264565, 264566, 64486, 264567	
	on .	4317445 (3929, 3930)	4107017 dbj BAA36294 -	ontains protein domain (PF01428) -		64488, 264510, 264760, 264768, 264486	
				N1-like Zinc finger			

996	94192058 (3931, 3932)		Contains protein domain (PF01027) - glycoprotein		22278999, 264092, 264259, 29331826,
		gil4929707/gb/AAD34114.1/AF15187 - (AF151877) CGI-119/Uncharactenzed protein family	9 Uncharacterized protein family		29331828, 29146498, 264595, 265011,
		protein [Homo sapiens]			264448, 18108354, 264288, 264684, 264766, 1 264685, 264686, 265022, 264601, 264692
					18108370, 18108377, 264555, 18108381.
					18108385, 264486, 264567
1961	87396123 (3933, 3934)	87396123 (3933, 3934) Novel Protein sim. GBank gij2957270 (AF044576) -	Contains protein domain (PF00388) - esterase		29331824, 265010, 265017, 264288,
		phospholipase C PLC210 [Caenorhabditis elegans]	Phosphatidylinositol-specific phospholipase C, X domain		21906764, 263981, 56526486
1868		88095641 (3935, 3936) Novel Protein sim. GBank gi[2564953 (AF030001) -	Contains protein domain (PF00008) - oncogene		35696286, 264905, 264509, 264906, 264907,
		unknown (Mus musculus)	EGF-like domain		264908, 264909, 264511, 264512, 265008,
					264910, 265009, 264594, 264757, 264758.
					264604, 264605, 264760, 264762, 264682,
					264764, 264685, 264766, 264767, 264689,
					264691, 264693, 264628, 264629, 35696423,
					35695855, 264631, 264632, 264634, 264635,
					264636, 264637, 18108380, 264564, 264565,
					264566, 264567
1969	84328529 (3937, 3938)	84328529 (3937, 3938) Novel Protein sim. GBank gi 2911274 (U20329) - spidroin 1		UNCLASSIFIED	22278995, 22278996, 35696052, 264906,
	_	(Nephila clavipes)			264908, 18108351, 264482
1970		Novel Protein sim. GBank gij4050087 (AF109907) - S164		UNCLASSIFIED	264908, 264288, 264766, 264636
		[Homo sapiens]			
1971	94843914 (3941, 3942)			collagen	264488, 264489, 22278998, 264259,
		gij134206 sp P09593 SANT_PLAFV - S-ANTIGEN			60432049, 66714117, 29331826, 60432289,
		PROTEIN PRECURSOR			29331827, 35696052, 264508, 264905,
					264509, 264906, 264907, 264908, 264909,
					264510, 264511, 264512, 264910, 264591,
					264592, 60432229, 60433356, 264595,
	•				264596, 264600, 264604, 264605, 264760,
					18108351, 264448, 264764, 264288, 264766.
					264768, 264769, 21906765, 33657023,
					264692, 18108370, 264629, 35696423,
					65274791, 35695855, 264632, 264635,
					264555, 264636, 264637, 264638, 264639.
					18108385, 60432113, 22279000, 264563,
27.0	12 CA 12 13 13 13 13 13 13 13 13 13 13 13 13 13	A STORE A Sell-Library Control of the Control of th	(C) 10 C		204304, 204303, 204300, 204400
7/6	יסיסיטייי (טשיט, טשיט)	orogenta (5545), 5544) Invover Frotein Siii. Gbaink gips 15025jaajja2450071.1j - [AB017546) homologois to the veest VGR163 gene [Miss	Contains protein domain (PPU1402) -		60432289 204633, 69331066, 30106101.
		musculus]	domain		265006 264511 285008 52644296 265018.
					265019, 264761, 264889, 21906768,
					21906769, 264691, 264693, 33657109,
					33657182, 264556, 52644332, 264558,
					60432113
1973	86395533 (3945, 3946)			UNCLASSIFIED	29331826, 264692, 35696423, 264631,
	_				264555, 264556, 264557, 264558, 264559
1974		80396629 (3947, 3948) Novel Protein sim. GBank gij3309543 (AF036382) - MLL Futu nibrines		UNCLASSIFIED	264682, 264764, 264563
			J		

UNCLASSIFIED 264488, 66714117, 29331826, 29331828, 56182435, 265006, 264757, 55812038, 265010, 265017, 264369, 55811957, 65274791, 35695855, 56182323, 60432113	UNCLASSIFIED 264259, 35696052, 265018, 265020, 265021, 33657109, 56526486		olfactor 60170831, 264566	UNCLASSIFIED 29331826, 29146498, 264905, 264907, 265007, 265009, 265010, 265018, 264686. 18108359, 21906768, 35695917, 265020, 60170615, 264693, 18108368, 18108370, 264631, 264635, 264556, 264558, 18108384, 22279000, 264565		\$SIFIED 22278995, 222789997, 22278999, 264259, 29331825, 29331828, 29146499, 294107, 264107, 264906, 264910, 264595, 21906754, 265010, 265017, 265018, 265019, 264448, 264288, 21906767, 33657023, 27468264, 18108370, 18108374, 18108375, 264630, 264631, 264635, 18108385, 87168518, 27279010, 284487, 284684	65274572, 29331825, 35696052, 33656970, 264909, 265008, 55811386, 264760, 264686, 244691, 27486264	
UNCLA	UNCLAS	потеорох (transcriptfactor	NCLA	ubiquitin	UNCLASSIFIED		Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type
949, 3950)	951, 3952)	94852664 (3953, 3954) Novel Protein sim. GBank gi[2499526]sp[Q07782]NASU_RAT - SODIUM/SULFATE COTRANSPORTER (NA(+)/SULFATE COTRANSPORTER)	87447645 (3955, 3956) Novel Protein sim. GBank gij103421/pir A33471 - Iranscription factor NTF1 - fruit fly (Drosophila melanogaster) (fragment)	87627709 (3957, 3958) Novel Protein sim. GBank gij2244815[embjCAB10238.1] - (297336) hypothelical protein [Arabidopsis thaliana]	86577059 (3959, 3960) Novel Protein sim. GBank gil4759290 ref NP_004642.1 pUSP1 - Ubiquitin carboxyl- terminal hydrolase, X-linked	87606974 (3961, 3962) Novel Protein sim. GBank gil4406693[gb AAD20060 - (AF131849) Unknown [Homo sapiens]	90995367 (3963, 3964) Novel Protein sim. GBank gi 5689523 db BAA83045.1 - (AB029016) KIAA1093 protein [Homo sapiens]	95098668 (3965, 3966) Novel Protein sim. GBank gij3417297 (AC002310) - Unknown gene product [Homo sapiens]
			1978 87447645 (39			1981 87606974 (39	1982 90995367 (39	1983 95098668 (39

84	85760989 (3967, 3968)	1984 85760989 (3967, 3968) Novel Protein sim. GBank gi 2896695 emb CAA17174.1 -		synthase	264688, 21906766, 55811957, 56994075,
		(ALUZ1897) radD14 [Mycobactenum tuberculosis]			200020, 200021, 22270999, 200022, 204209, 100001800 044209, 100001800 044600
					18108370, 264908, 264629, 55811576.
					35695855, 265006, 265007, 264591,
					21906754, 33657084, 265010, 265017,
					265019, 264288
1985		85636897 (3969, 3970) Novel Protein sim. GBank		glycoprotein	264760, 264288, 263978, 55811576, 264637,
_		gij5712131 gb AAD47379.1 AF12049 - (AF120499) DEM1 protein [Homo sapiens]			56182323, 18108385, 264564
986	80200507 (3971, 3972)			UNCLASSIFIED	264488, 264629
1987	87011117 (3973, 3974)	87011117 (3973, 3974) Novel Protein sim. GBank	Contains protein domain (PF00069) -		22278999, 29331830, 265007, 265018,
		gi 4868443 gb AAD31319.1 AF14457 - (AF144573) Mx-	Eukaryotic protein kinase domain		21906768, 33657023, 264692, 264693,
		interacting protein kinase PKM [Mesocricetus auratus]			18108377, 264635, 60170394, 22279002
1988	94122108 (3975, 3976)			UNCLASSIFIED	264905, 264906, 264907, 264908, 264909,
					264910, 264591, 264593, 264758, 264764,
					264686, 264768, 265021, 264692, 264628,
					264629, 35695855, 264630, 264635, 264636,
					264637, 264638, 264639, 264483
1989	91225225 (3977, 3978)	91225225 (3977, 3978) Novel Protein sim. GBank gi[2801701 (AF042379) - spindle		tubulin	60432049, 60432289, 52644045, 56182435.
		pole body protein spc97 homolog GCP2 [Homo sapiens]			264112, 265007, 33657402, 52644229,
					21906765, 21906768, 21906769, 55811957.
					33657023, 263967, 33657109, 18108370,
					22279000, 22279002
1990	85699888 (3979, 3980)	85699888 (3979, 3980) Novel Protein sim. GBank gi 5701727 db BAA83074.1 -			264508, 264757, 264764, 18108381
		(AB024729) alpha-1,3-D-mannoside beta-1,4-N-			
		acetylglucosaminyltransferase IV-homologue [Homo			
1991	95353114 (3981 3982)	95353114 (3981 3982) Novel Protein sim GBank gild240287 IdbilBAA74922 11-	Contains protein domain (PF01602) - glycoprotein	alvcoarotein	18108394, 56182575, 22278994, 35696286,
		(AB020706) KIAA0899 protein [Homo sapiens]	Adaptin N terminal region		56994075, 22278997, 22278999, 29331822,
					29331824, 29331825, 60432289, 29331828,
					264508, 264906, 264907, 264908, 56182435,
					264510, 265007, 21906754, 33109954,
					87168474, 265017, 265018, 265019, 264762.
					18108351, 264763, 264683, 264369, 264288,
					264685, 264766, 264687, 264769, 21906765,
					21906768, 21906769, 55811957, 265020,
					60431528, 263974, 18108379, 35695855,
					264555, 264557, 264639, 83373044,
					18108384, 87168518, 60432113, 22279000,
1					22279002, 264564, 264486
1882	95317232 (3983, 3984)	95317232 (3983, 3984) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73,	Contains protein domain (Pruduse) - struct	Siruct	28331827, 2849U6, 2849U7, 2849U9, 283UU7.
		contains large complex repeat CR 73 [Kaposi's sarcoma-	Zinc finger, C2H2 type		264603, 264765, 264685, 264766, 21906766,
		associated herpesvirus]			264628, 264633, 264636, 18108363, 56526486, 264566, 264567
1993	-	80054763 (3985, 3986) Novel Protein sim. GBank gi 2565091 (U80761) - CTG26		UNCLASSIFIED	264592, 35696423
		alternate open reading frame [Homo sapiens]			

264486, 22278997, 22278998, 22278999, 29331822, 29331822, 29331822, 29331826, 29331826, 29331826, 29331826, 29331826, 29331827, 29331828, 264905, 264907, 264908, 265006, 6043229, 33657402, 6043336, 264757, 60433438, 264758, 33109954, 265011, 265017, 265018, 265019, 264684, 264369, 264685, 264686, 21906769, 21906765, 21906769, 18108362, 264693, 65274620, 18108370, 264635, 264556, 26457, 56182323, 83373044, 56528486, 22279000, 22279002, 264564	264488, 18108396, 22278994, 56994075, 22278996, 35696286, 22278997, 22278999, 264259, 29147620, 56182181, 29331824, 6043289, 2934826, 29331827, 35696052, 29146499, 264905, 264907, 66712502, 56182435, 265007, 265008, 265009, 56182435, 265008, 265009, 265010, 265011, 265017, 265018, 265019, 264761, 264761, 264764, 264686, 264768, 264769, 264288, 264766, 294686, 21906765, 21906766, 21906767, 29148627, 21906765, 21906769, 29148629, 29148784, 265020, 265021, 264690, 18108377, 36996423, 55811576, 65274791, 3569555, 264631, 20431850, 264555, 264556, 20431850, 264557, 264558, 264566, 20431850, 264557, 264558, 264565, 264431, 264567, 20438518, 264567, 20431871, 264558, 264556, 264531, 264567	264564	264908, 264909, 264592, 264593	264113, 264685, 264555, 264567
rnapolymerase	UNCLA	UNCLASSIFIED	phosphatase	UNCLASSIFIED
Contains protein domain (PF00856) - mapolymerase SET domain			Contains protein domain (PF00023) - phosphatase Ank repeat	
1994 94329114 (3987, 3988) Novel Protein sim. GBank gi 5630077 gb AAD45822.1 AC00601 - (AC006017) similar to AAC51735 (PID:g2358287) Homo sapiens	95414353 (3989, 3990) Novel Protein sim. GBank gil4827040[ref[NP_005110.1 pTRAP - thyroid hormone receptor-associated protein, 150 kDa subunit	80254186 (3991, 3992) Novel Protein sim. GBank gi 791146 emb CAA60020 - (X86028) extensin-like protein [Vigna unguiculata]	gij2642034 (AF034547) - protein i binding subunit [Ovis aries]	85262704 (3995, 3996) Novel Protein sim. GBank gil4589634 dbj BAA76839.1 - (AB023212) KIAA0995 protein [Homo sapiens]
94329114 (3987, 3988)	95414353 (3989, 3990)		87028423 (3993, 3994)	85262704 (3995, 3996)
4994	330			1998

1899	94324903 (3997, 3998)	1999 94324903 (3997, 3998) Novel Protein sim. GBank gij5225312 gb AAD40846.1 AF07244 - (AF072441) calcineurin binding protein cabin 1 [Homo sapiens]	Contains protein domain (PF00515) - UNCLASSIFIED TPR Domain	UNCLASSIFIED	18108394, 18108397, 35696286, 60424269, 29331827, 29331828, 35696526, 265006, 264512, 55811386, 265010, 265018, 265019, 55811150, 18108351, 264763, 264682, 264369, 264682, 264691, 33657023, 264686, 56181562, 265020, 27486264, 18108370, 18108379, 35695855, 264634, 264555, 264557, 56182323, 18108382, 264559, 83373044, 60432113, 22279000, 264563, 264564,
		Novel Protein sim. GBank gij1723232[spjQ10155]YATA_SCHPO - HYPOTHETICAL 90.6 KD PROTEIN C1D4.10 IN CHROMOSOME I		UNCLASSIFIED	52646365, 52646842, 22278894, 22278895, 22278896, 35686286, 22278897, 22278898, 326278998, 32278998, 32278998, 32278998, 32278998, 32278998, 32278998, 3237829, 22278998, 3237829, 22278998, 3237828, 3237828, 3264106, 29331820, 22644045, 265003, 60170831, 264592, 264593, 32657402, 60433438, 21906765, 21906766, 21906767, 21906768, 21906769, 35695917, 265020, 265021, 60170815, 52644150, 33657023, 65274620, 52644332, 5486262, 27486264, 35695763, 35695853, 35695855, 264631, 52644332, 56182323, 60170394, 83373044, 56526486, 22279002, 264566, 264567
	95072534 (4001, 4002)	95072534 (4001, 4002) Novel Protein sim. GBank gi 107560 pir B38637 - Ras inhibitor (clone JC265) - human (fragment)		UNCLASSIFIED	264769, 52644229, 65274572, 21906768, 22278996, 3569586, 35695917, 265020, 22278999, 284534, 26490, 264259, 264692, 66432289, 33657109, 35696052, 284508, 35696423, 65274791, 35695655, 264510, 264511, 264512, 265009, 284634, 264536, 264556, 264558, 264559, 264559, 264559, 264601, 264601, 264601, 264601, 264601, 264601, 264288, 264765
	80236368 (4003, 4004)	80236368 (4003, 4004) Novel Protein sim. GBank gil729433 sp P38657 ER60_BOVIN - PROBABLE PROTEIN Thioredoxin DISULFIDE ISOMERASE ER-60 PRECURSOR (ERP60) (58 KD MICROSOMAL PROTEIN) (P58)	olein domain (PF00085) -	isomerase	264907, 265006, 264910, 264603, 264692, 264629, 18108374, 264556, 264557
2003	80074449 (4005, 4006)	80074449 (4005, 4006) Novel Protein sim. GBank gi[86388 pir A27040 - neurofilament triplet M protein - chicken (fragment)		UNCLASSIFIED	264905, 264906, 264908, 264910, 264596, 265017, 18108351, 264692, 264629, 264634, 264565

200	95317318 (4007, 4008)	2004 95317318 (4007, 4008) Novel Protein sim. GBank gij484249lemblCAB43230.1 - (AL049996) hypothetical protein [Homo sapiens]	Contains protein domain (PF00076) - dna_ma_bind RNA recognition molif. (a.k.a. RRM, RBD, or RNP domain)		52645156, 52646842, 52646365, 56182575, 52278994, 22278994, 22278996, 35696266, 22278996, 35696266, 22278996, 35696266, 22278999, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22331824, 22331826, 2331826, 2331826, 2331826, 2331826, 23357402, 52646317, 21906754, 33657084, 5264429, 2646317, 265017, 265018, 265019, 264763, 264683, 265020, 265021, 265022, 52644150, 23657023, 52645129, 23657109, 33657182, 27486261, 27486262, 33657189, 27486265, 35695673, 18108376, 18108376, 18108377, 266458, 18108385, 56526486, 87168518, 264558, 18108385, 56526486, 87168518,
2005	87400864 (4009, 4010)	87400864 (4009, 4010) Novel Protein sim. GBank gil3879501 lemb[CA487795] - (Z47812) similar to ubiquitin carboxyl-terminal hydrolase; cDNA EST EMBL:D33366 comes from this gene; cDNA EST EMBL:D33965 comes from this gene; cDNA EST EMBL:D33822 comes from this gene; cDNA EST EMBL:D34547 comes from this gene; cDNA EST		ubiquitin	264488, 264906
	95351177 (4011, 4012)	95351177 (40†1, 40†2) Novel Protein sim. GBank gi 4106673 emb CAA22613 - (AL035064) queuine tma-ribosyltransferase [Schizosaccharomyces pombe]	Contains protein domain (PF01702) - UNCLASSIFIED Queuine tRNA-ribosyltransferase		56182575, 56181686, 22278995, 22278996, 22278997, 22278998, 22278999, 264259, 56182181, 60444269, 66714117, 35696052, 264906, 66712502, 264909, 264510, 60433356, 8568842, 265010, 265018, 265019, 264682, 26448, 264288, 264768, 29148627, 21906769, 29148784, 35693917, 60170615, 264691, 33657109, 55810764, 55811576, 35695855, 87168518, 60432113, 264563, 264482
2007	94325556 (4013, 4014)	94325556 (4013, 4014) Novel Protein sim. GBank gi 2662161 db BAA23712 - (AB007900) HH0452 CDNA clone for KIAA0440 has a 438-bp insertion at position 1711 of the sequence of KIAA0440. [Homo sapiens]		UNCLASSIFIED	264488, 263994, 35696052, 264508, 264905, 264906, 264907, 264908, 264909, 265007, 264908, 264909, 265007, 264908, 264909, 265007, 264900, 264768, 264768, 264768, 264768, 264685, 264677, 264678, 264691, 264693, 264629, 26534791, 35695855, 264631, 264632, 264634, 264635, 264637, 264638,
2008	85084428 (4015, 4016)	85084428 (4015, 4016) Novel Protein sim. GBank gi]1550783 emb CAA69257 - (Y07960) homeodomain protein [Mus musculus]	Contains protein domain (PF00046) - homeobox Homeobox domain		264909, 264768, 35695855

 600	85749240 (4017, 4018)	2009 85749240 (4017, 4018) Novel Protein sim. GBank gij3882305 dbj BAA34512.1 - (AB018335) KIAA0792 protein [Homo sapiens]		UNCLASSIFIED	22278999, 264259, 264910, 264591, 265017, 264681, 264683, 21906768, 264691,	
	_				3353/162, 3353/349, 264531, 8/168318, 264404, 22279002, 264563	
2010		95422458 (4019, 4020) Novel Protein sim. GBank gij5262629jemb CAB45753.1 - (AL080164) hypothetical protein [Homo sapiens]	Contains protein domain (PF00057) - eph Low-density lipoprotein receptor	hqa	52644507, 52645156, 52646365, 52646842, 18108397, 65274572, 22278994, 56994075,	
			domain class A		35696286, 22278996, 22278997, 22278999,	
					29331825, 29331826, 29331827, 29331828, 29331825, 29331826, 29331827, 29331828,	
					264511, 265007, 264512, 265008, 265009,	
					60432229, 60433356, 21906754, 52646317,	
					33109954, 52644296, 87168474, 87168559, 265017, 265018, 264685	
					264687, 52644229, 264689, 21906765,	
					21906766, 21906767, 21906768, 35695917,	
					265020, 52644150, 264691, 264692,	
					33657023, 263967, 52645129, 35695763,	
					18108376, 35696423, 65274791, 35695855,	
					264631, 264634, 60431850, 264637, 264638,	
					52644332, 60170394, 18108385, 87168518,	
					22279002, 264564, 264565, 264566, 264567	_
2011		94328149 (4021, 4022) Novel Protein sim. GBank gi(3347953 (AF076183) -		UNCLASSIFIED	56182575, 56994075, 22278999, 264259.	_
		cytosolic sorting protein PACS-18 [Rattus norvegicus]			29331624, 29331626, 29331627, 29331626.	
					JOSEPH ST. SEASON DEFO. DEFO. SESSION CONTROL	
					2650U7, 2650U8, 265U10, 265U11, 265U17,	
					203019, 204001, 204440, 204003, 204309,	
					204486, 204885, 204786, 204887, 21908785, 31008787, 310087888, 310087888, 310087888, 310087888, 310087888, 3100888888, 310088888, 31008888, 31008888, 310088888, 31008888, 31008888, 31008	
					21300101, 21300100, 21300103, 203020, 2650200, 265020, 265020, 265020, 265020, 2650200, 2650200, 2650200, 2650200, 2650200, 2650200, 2650200, 2650200, 2650200, 26502000, 2650200, 2650200, 2650200, 2650200, 2650200, 2650200, 26502000, 26502000, 26502000, 26502000, 26502000, 26502000, 26502000, 265020000, 265020000, 265020000, 265020000000000000000000000000000000000	
					33657109, 264629, 264557, 264559,	
					83373044, 87168518, 60432113, 22279002	
2012	87772137 (4023, 4024)	87772137 (4023, 4024) Novel Protein sim. GBank gij1086678 (U41020) - coded for	Contains protein domain (PF00409) - UNCLASSIFIED	UNCLASSIFIED	264259, 29331822, 29331824, 29331825,	
		by C. elegans cDNA yk100g4.5; coded for by C. elegans	Kinesin light chain repeat		29146498, 87168559, 265019, 264682, 254388, 254688, 24008764, 255030, 255033	
		belix loop belix lancing along transminion factor			204288, 204880, 21988/84, 283820, 283822, 33657833 364683 33657488 65811576	
		[Caenorhabditis elegans]			264632, 264558, 56182323, 264639,	
		•			18108383, 18108384, 18108388, 22279000,	
3					22279002, 264567	
5102	94843842 (4025, 4026)		Contains protein domain (PF00096) - dna_rna_bind	dua_ma_bind	18108398, 264908, 265007, 265010, 265018.	_
		gi 4507985 ref NP_003427.1 pZNF1 - zinc finger protein 135 Zinc finger, C2H2 type (clone pHZ-17)	Zinc finger, C2H2 type		265019, 264689, 21906767, 265020, 264692	
2014		87347940 (4027, 4028) Novel Protein sim. GBank		UNCLASSIFIED	264488, 29331826, 264907, 264636, 264555.	
		gi(127720)sp P20938 MYPO_HETFR - MYELIN PO PROTEIN PRECURSOR			264639, 264558	
1		יו אסובווען ואבסטומסוי				

2015	88094922 (4029, 4030)	2015 88094922 (4029, 4030) Novel Protein sim. GBank gil81286lpir S22697 - extensin -		UNCLASSIFIED	56182575 35696286 264259 35696052
		Volvox carteri (fragment)			264508, 264906, 264907, 264510, 264512,
					87168474, 265010, 264681, 264288, 264689,
					264628, 35696423, 35695855, 264639,
					264563, 264564
2016		85298641 (4031, 4032) Novel Protein sim. GBank gi 285046 pir S26413 - t-complex		struct	264102, 264508, 264110, 265009, 33109954,
					Z1905/05, Z030Z1, 3303/109, Z/480Z0Z, 263972, 18108374, 263976, 264555, 264564
2017				UNCLASSIFIED	264685, 264636
2018		79637067 (4035, 4036) Novel Protein sim. GBank gij124735 sp P18175 INVO_PIG-			264693
3	_	INVOLUCIAIN			
<u> </u>		87787900 (4037, 4038) Novel Protein sim. GBank gi[2143910]prr 568216 - Inhosophalase-1 nivcogen-hinding (GL)-chain - rat		phosphatase	264107, 264110, 264112, 265017, 263976
2020		94674476 (4039 4040) Novel Protein sim GBank 012078483 (1143200) - antifreeze		INCLASSIEIED	264250 264508 264501 265018 264682
_		glycopeptide AFGP polyprotein precursor [Boreogadus saida]			264288, 264688, 22279002
203	Т	86718818 (4041 4042) Navel Protein eim CBack			66004076 264602 22100064 21006764
		AISSECTION SILL COMING ON EL CHONGATION			3003340/3, 204333, 33103334, 21306/34,
		FACTOR G, MITOCHONDRIAL PRECURSOR (MEF-G)			87168518
2022		95295665 (4043, 4044) Novel Protein sim, GBank gil4218005 (AC006135) - outstive			264757 264767 60170615 18108385
		vicilin storage protein (globulin-like) [Arabidopsis thaliana]			
2023	87722976 (4045, 4046)	2023 87722976 (4045, 4046) Novel Protein sim. GBank	Contains protein domain (PF00442) - ubiquitin	ubiquitin	18108394, 22278999, 264259, 264905.
		gi 5410230 gb AAD42992.1 AF07334 - (AF073344) ubiquitin Ubiquitin carboxyl-terminal	Ubiquitin carboxyl-terminal		264906, 264908, 264595, 264762, 264769.
		specific protease 3 [Homo sapiens]	hydrolases family 2		264634, 264636, 87168518, 60432113,
	~				22279000, 264482, 264565
2024	87896443 (4047, 4048)	-			60433438, 265017, 264686, 264692, 264693,
2025		87858863 (4049 4050) Novel Protein sim GBank	Contains protein domain (PE00637) - HINCLASSIFIED	INCI ASSIEIED	22278997 264509 264906 264909
		1 AC00701 - (AC007018)	7-fold repeat in Clathrin and VPS		55812038, 265017, 265021, 265022,
			•		60170615, 264556
2026		94122114 (4051, 4052) Novel Protein sim. GBank gi 1655699 emb CAA69032 -		UNCLASSIFIED	56994075, 60432049, 264508, 66712502,
		(Y07752) pherophorin-S [Volvox carteri]			264112, 60170831, 87168559, 264288,
					264688, 264689, 21906766, 33657109,
					18108370, 264638, 18108385, 60432113,
					22279000, 22279002, 264564, 264566,
7007					26436/
7707	80249001 (4053, 4054)			UNCLASSIFIED	263978, 264634, 264486

8028 5028	94699884 (4055, 4056	2028 94699884 (4055, 4056) Novel Protein sim. GBank gij1330345 (U58755) - coded for by C. elegans cDNA yk34b1.5; coded for by C. elegans cDNA yk13h10.5; coded for by C. elegans cDNA yk46e8.5; coded for by C. elegans cDNA yk46685; coded for by C. elegans cDNA yk46685.5; coded for by C. elegans cDNA yk46e8.5. yk46e8	11330345 (U58755) - coded for Contains protein domain (PF00632) - ubiquitin 5; coded for by C. elegans HECT-domain (ubiquitin-by C. elegans cDNA yk46e8.5; transferase). A yk46d5.5; coded for by C. elegans cDNA defects of the coded for by C. elegans cDNA	ubiquitin	52644507, 52645156, 52646842, 56182575, 56994075, 35696286, 22278997, 22278998, 22278999, 62278999, 22278998, 22278999, 60432049, 264259, 52645080, 29331822, 29331824, 66714117, 29331825, 25331828, 35696052, 264906, 264908, 29331830, 52644045, 56182435, 265009, 50442229, 33657402, 264509, 264757, 5264100, 264410, 564410, 56544, 52646317, 52644266, 55644, 5264431, 5264501, 5264426, 5264431, 5264501, 5264426, 5264431, 5264501, 5264426, 5264431, 5264501, 5264426, 5264431, 5264501, 5264426, 5264431, 526441, 526441, 526441, 526441, 52641, 52641, 52641, 52641, 52641, 52641, 52641, 52641, 52641, 52641, 52641, 52641
					265019, 265011, 67160359, 265011, 265010, 265019, 265019, 265019, 265019, 264766, 21906766, 21906767, 21906767, 21906767, 21906767, 21906767, 21906769, 5811957, 25695917, 265020, 265021, 265022, 25644150, 33657023, 65274620, 33657109, 52645129, 18108368, 27486261, 27486262, 27486264, 27486265, 35695763, 264629, 2646335, 2646335, 2646335, 2646336, 2646335, 2646336, 26463
5029	95362032 (4057, 4058)	2029 95362032 (4057, 4058) Novel Protein sim. GBank gil3599940 (AF017368) - faciogenital dysplasia protein 2 [Mus musculus]	Contains protein domain (PF00621) - UNCLASSIFIED RhoGEF domain	UNCLASSIFIED	265009, 264585, 85658542, 264555, 264556, 264555, 264556, 264557, 264557, 264558, 264559, 83373044
2030		91213734 (4059, 4060) Novet Protein sim. GBank gil5630080[gb]AAD45825.1]AC00489 - (AC004890) similar to HUB1; similar to BAA24380 (PID:g2789430) [Homo sapiens]	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	18108394, 56994075, 22278997, 22278999. 264259, 29331822, 29331824, 29331825, 66714117, 60432289, 29331826, 264108, 66712502, 264828, 265009, 265018, 265019
					264681, 264682, 264684, 264685, 56181562. 264689, 21906769, 265022, 264692, 264693. 264628, 18108370, 18108374, 264634, 264636, 264556, 18108385, 87168518, 22278002, 284565
2031					264591, 55811957, 18108365, 264557, 264558, 18108382, 18108384
2032		91232607 (4063, 4064) Novel Protein sim. GBank gi 5689491 dbj BAA83029.1 - (AB029000) KIAA1077 protein [Homo sapiens]	Contains protein domain (PF00884) - hydrolase Sulfatase	hydrolase	65274572, 35696286, 29331824, 264908, 265009, 264593, 265018, 264288, 264686, 284769, 21906766, 21906767, 29148627, 264628, 35696423, 264634, 264556, 18108381, 60170394, 264559, 83373044, 18108381, 574484
2033		2033 95000809 (4065, 4066) Novel Protein sim. GBank gi[2494828]sp[064686]CAG7_RAT - ALPHA-N-ACETYLGALACTOSAMINIDE ALPHA-2,6-SIALYLTRANSFERASE (ST6GALNACIII) (STY)		synthase	56181562, 264628, 264632, 264555, 264556
2034		91232529 (4067, 4068) Novel Protein sim. GBank gil4826984 ref NP_005147.1 pROD1 - UNKNOWN	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif, (a.k.a. RRM, RBD, or RNP domain)	dna_ma_bind	22278996, 22278999, 264907, 29331830, 265008, 265018, 264681, 264682, 264684, 21906767, 21906768, 21906769, 33657109, 83373044, 56526486

2035	2035 83553451 (4069, 4070)				
2036	2036 87115833 (4071, 4072)				29331827, 29331828, 264682, 264369,
2037		94324833 (4073, 4074) Novel Profein sim GBank All 2734081 (46000195)			29148627, 60432113
		to covered thinding proteins (Capacitation also and		UNCLASSIFIED	65274572, 22278995, 22278996, 56994075,
_		seronation of the continuous property of the continuous elegans!			35696286, 22278997, 22278998, 22278999,
					264259, 29331824, 60432289, 29331826,
	_		•		29331828, 35696052, 264907, 29331830,
					66712502, 56182435, 265008, 265009,
					60170831, 264594, 55812038, 33109954,
_					21906754, 87168559, 265017, 265018,
					265019, 264762, 264369, 264288, 21906765,
				4	21906767, 21906768, 21906769, 55811957.
					35695917, 265020, 265021, 265022,
					52644150, 33657023, 33657109, 33657182,
					35695763, 35695855, 264632, 264634,
					264636, 56182323, 83373044, 60432113,
3030	_				22279000, 22279002, 264563
2000		32422304 (4073, 4070) Novel Protein Sim. GBank gil3880625[emb[CAB07858] -	Contains protein domain (PF01412) - UNCLASSIFIED	UNCLASSIFIED	22278995, 22278996, 56994075, 264259,
		(293/85) predicted using Genetinder; similar to RNA	Putative GTP-ase activating protein		29331824, 35696052, 264905, 264906,
		recognition motif (aka RRM, RBD, or RNP domain); cDNA	for Arf		52644045, 265007, 265009, 87168559,
		EST EMBL: T01682 comes from this gene; cDNA EST			265017, 18108351, 264448, 264369, 264766.
		EMBL:M75823 comes from this gene; cDNA EST			264767, 264686, 18108358, 21906765,
		EMBL:D27559 comes from this ge			21906769, 52644150, 33657023, 264692,
					18108362, 33657109, 27486262, 18108370,
					18108374, 18108379, 35696423, 65274791,
					264632, 264636, 18108383, 83373044,
					18108385, 87168518, 22279000, 22279002,
0000	2000 1000 0000 1000				264563, 264564, 264566
5023	02214020 (4077, 4070)	2038 93314020 (4077, 4076) Novel Protein Sim. GBank gi 2224653 db BAA20813 -	Contains protein domain (PF00097) - UNCLASSIFIED	UNCLASSIFIED	22278997, 264259, 29331822, 264905,
		(Abutasa4) NiAAusab [Homo sapiens]	Zinc finger, C3HC4 type (RING		264906, 264907, 264908, 264909, 264510,
			finger)		265009, 264910, 264593, 264758, 265011,
					265018, 264762, 264288, 264766, 264768,
					264769, 21906766, 33657023, 264692,
					264693, 33657109, 35696423, 264631,
					264632, 264634, 264635, 264636, 264637,
9	0207 CEOTY ETT 00000				264639, 87168518, 264486
	_			UNCLASSIFIED	264592
4504				rnapolymerase	264488, 22278998, 35696052, 264905,
		gilzoubzajspir / 0/00jrPA2_MOUSE - DNA-DIRECTED			264907, 264908, 264910, 265018, 264605,
		ANA POLYMERASE 1135 KD POLYPEPTIDE (RNA			265019, 18108351, 264766, 264769,
		POLIMERASE I SUBUNIT 2) (RPA135)			21906766, 265021, 265022, 264692,
					33657109, 264628, 264629, 35696423,
					35695855, 264637, 264638, 264563, 264564,
					264565, 264567

2042	95307447 (4083, 40	384) Novel Pi	2042 95307447 (4083, 4084) Novel Protein sim. GBank gil4405590 gb AAD20040 -	Contains protein domain (PF00568) - UNCLASSIFIED	UNCLASSIFIED	60424179, 35696286, 264259, 29331826,
		(AF131)	(AF131766) Similar to Ena-VASP like protein [Homo	WH1 domain		35696052, 29331828, 264508, 264509,
		[sapiens]				264907, 264909, 264510, 264511, 265009,
_						264910, 264591, 60433356, 264595, 265017,
_						265019, 264681, 264764, 264369, 264765,
						264684, 264288, 264766, 264688, 52644229,
						264769, 21906765, 35695917, 264535,
						52644150, 264691, 264692, 18108365,
						27486261, 27486262, 27486265, 18108374,
						35696423, 65274791, 35695855, 264555,
						264558, 60170394, 18108385, 264404,
	-					22279000, 22279002, 264482, 264563,
	_	-				264564, 264566
2043	94328076 (4085, 4086) Novel Protein sim. GBank	386) Novel P		Contains protein domain (PF00122) - transport	transport	264488, 52644507, 52646365, 56994075,
	***	gi 5052£	gij5052554[gb]AAD38607.1[AF14563 - (AF145632)	E1-E2 ATPase		22278997, 22278999, 20281171, 264259,
		BCUNA.	GH06032 [Drosophila melanogaster]			29331822, 29331824, 66714117, 29331826,
						29331828, 33656970, 29146498, 264509,
						264908, 52644045, 56182435, 265006,
						33657402, 21906754, 52644296, 87168559,
						265017, 265018, 265019, 264681, 264288.
						264766, 264685, 264686, 21906766,
						21906767, 21906768, 21906769, 265020,
						265021, 60170615, 264691, 33657023,
						264693, 65274620, 33657109, 33657182,
						27486261, 27486262, 33657349, 35695763,
				٠		18108374, 55811576, 35695855, 18108380,
						18108381, 60170394, 56182323, 264558.
						83373044, 18108385, 56526486, 87168518.
						60432113, 22279000, 264567
2044	87106927 (4087, 4088) Novel Protein sim. GBank of	383) Novel Pr	rotein sim. GBank gil2246532 (U93872) - ORF 73		struct	264093 29331827 264905 66712502
:		contains	contains large complex repeat CR 73 IKaposi's sarcoma-			264592, 264689, 21906765, 21906769,
		associat	associated herpesvirus]			265020, 264692, 264482, 264566
2045	_	(060				264692
2046	_	192) Novel Pi	87320849 (4091, 4092) Novel Protein sim. GBank gil4406698 gb AAD20062 -			264259, 264906, 264683, 22279002
7047	1	Moved Dy	R4578801 (4003 4004) Novel Protein eim CBank gild 101720 (AEOOG468)			22278000 20147620 20231824 20146408
<u> </u>	_	lymphoc	wmphocyte specific formin related protein fMus musculus			264508 265007 265008 265019 264605
						264681 20148627 20148620 266021
						204081, 23146027, 23146025, 203021. 33657023 48108366 33667100 33667182
						3303/023, 10100303, 3303/109, 3303/102,
						18108377, 264556, 264638, 264559,
						18108388
2048		196)			UNCLASSIFIED	264909
2049		398) Novel Pi	88094690 (4097, 4098) Novel Protein sim. GBank gil4589656 dbj BAA76850.1 -		UNCLASSIFIED	264488, 264259, 29331824, 29331828,
_		(AB0232	(AB023223) KIAA1006 protein [Homo sapiens]			35696052, 264906, 264907, 264908, 264909,
						264910, 264603, 264763, 21906767,
						21906768, 264629, 264634, 264637,
						22279002, 264564, 264565, 264566, 264567

2061	95362204 (4121, 4122)	95362204 (4121, 4122) Novel Protein sim. GBank	Contains protein domain (PF00069) - kinase	kinase	22278997, 22278999, 264259, 29331822.
		gi 2496947 sp Q09298 YQO9_CAEEL - HYPOTHETICAL	Eukaryotic protein kinase domain		29331824, 29331826, 29331827, 35696052,
		141.2 KD PROTEIN EEED8.9 IN CHROMOSOME II			29331828, 264906, 66712502, 29331830,
					264908. 264909. 264112, 264511, 265007.
					265009, 264910, 264591, 33657402,
					21906754, 85658542, 265017, 265019,
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			•		264685, 264687, 18108358, 264689,
					21906765, 21906767, 21906768, 21906769,
					265020, 265022, 264691, 33657023,
					33657109, 20281149, 18108379, 35695855,
					264634, 264556, 264557, 264558, 18108382,
					264559, 83373044, 18108384, 56526486,
					60432113
2062			Contains protein domain (PF00023) - struct	struct	264905, 264628, 264907, 264629, 264908,
		gil4502091 refine _001139.1 pANK2 - ankyrin 2, neuronal	Ank repeat		264909, 18108374, 263978, 35695855,
					264512, 264635, 60431850, 264636, 264760,
					264563, 18108351, 264762, 264565, 264764,
					264487, 264766
2063		87601272 (4125, 4126) Novel Protein sim. GBank gil4589562 dbj BAA76803.1 -	Contains protein domain (PF00617) - oncogene	oncogene	22278994, 22278999, 264259, 29331827,
		(AB023176) KIAA0959 protein [Homo sapiens]	RasGEF domain		264906, 264909, 52644045, 264686,
					21906767, 55811957, 264692, 18108365,
					263972, 55811576, 18108384, 22279002,
					264482, 264563, 264564, 264484
2064		95317253 (4127, 4128) Novel Protein sim. GBank gi 1754515 dbj BAA13413.1 -		hydrolase	264488, 52646365, 56994075, 35696286,
		(D87515) aminopeptidase-B [Rattus norvegicus]			22278997, 22278998, 264259, 29331826,
					60432289, 29331827, 29331828, 35696052,
					264509, 265007, 265008, 60432229,
					60433438, 21906754, 265010, 265011,
					87168559, 265017, 265018, 264761,
					18108351, 264682, 264369, 264288,
					52644229, 21906765, 21906767, 21906768.
					35695917, 33657109, 18108368, 18108374,
					35696423, 35695855, 52644332, 264559.
					60432113, 22279000, 22279002, 264566.
					264486
2065		95092238 (4129, 4130) Novel Protein sim. GBank		kinase	264569, 18108394, 56182181, 60432289,
		gi 2507144 sp Q04205 TENS_CHICK - TENSIN			29331826, 264905, 264906, 264908,
					60431735, 60433356, 55811386, 85658542,
					265018, 55811150, 264681, 264766, 264692,
					60431528, 263974, 55810764, 35695855,
					264631, 264634, 264635, 60431850, 264557,
					83373044, 18108388, 22279000, 22279002
2066		85793402 (4131, 4132) Novel Protein sim. GBank gil160171 (M58295) -	Contains protein domain (PF00096) - UNCLASSIFIED	UNCLASSIFIED	56182575, 264259, 264906, 264764, 264288,
		crcumsporozoite protein [Plasmodium yoelii]	Zinc tinger, CZHZ type		56182323, 264567

35696286, 22278997, 22278998, 60432049, 264259, 60432289, 60433438, 264682, 264448, 264369, 264288, 18108355, 21908765, 21906768, 265022, 33657109, 35696423, 35695855, 264558, 264404, 264563, 264486	264687	60424179, 56182575, 22278995, 22278996, 56994075, 264259, 29331822, 29331824, 29331825, 29331824, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 264331825, 265607, 265609, 264356, 2656017, 265018, 264604, 265019, 264418, 264764, 264766, 21966765, 21966767, 21966767, 21966769, 263976, 264555, 264557, 56182323, 83373044, 87168518, 60432113, 22279000, 22279002	35695917, 264905, 264628, 264908, 264638	18108398, 265006, 265007, 265008, 265009, 264594, 265010, 265011, 18108351, 18108354, 18108368, 18108365, 18108388, 264634, 18108385, 18108388, 18108391	22278995, 35696286, 22278997, 22278998, 22278999, 264490, 60432049, 264259, 29331824, 29331826, 3569652, 265008, 33657402, 21906754, 265011, 265019, 18108351, 264682, 264369, 21906765, 21906766, 21906767, 21906768, 264692, 35696423, 264555, 264556, 264558, 22279000	264556	29331822, 264909, 264511, 265009, 264594, 264596, 264596, 265011, 265011, 265017, 265018, 265019, 265018, 265020, 265021, 18108370, 264632, 83373044, 264567
	UNCLASSIFIED	transcriptfactor	UNCLASSIFIED			UNCLASSIFIED	
		Contains protein domain (PF00023) - transcriptfactor Ank repeat		Contains protein domain (PF00568) - WH1 domain	Contains protein domain (PF00184) - Neurohypophysial hormones, C- terminal Domain		
		94319177 (4137, 4138) Novel Protein sim. GBank gij3152662 (AF064604) - KE03 protein [Homo sapiens]	85791380 (4139, 4140) Novel Protein sim. GBank gi 5712131 gb AAD47379.1 AF12049 - (AF120499) DEM1 protein [Homo sapiens]	86946116 (4141, 4142) Novel Protein sim. GBank gil3551531 dbj BAA33016 - (AB017437) avena [Gallus gallus]		27925664 (4145, 4146) Novel Protein sim. GBank gi 1504026 dbj BAA13212 - (D86976) similar to C.elegans protein (237093) [Homo sapiens]	2074 94324767 (4147, 4148) Novel Protein sim. GBank gi 4240317 dbj BAA74937.1 - (AB020721) KIAA0914 protein [Homo sapiens]
2067 95303892 (4133, 4134)			85791380 (4139, 4140)		91718429 (4143, 4144) ,	27925664 (4145, 4146)	94324767 (4147, 4148)
2067	2068	2069	2070	2071	2072	2073	2074

18108394, 22278994, 22278996, 35696286, 22278998, 22278998, 264259, 29331822, 29331825, 29331827, 35696052, 29331828, 264908, 264908, 264910, 265009, 33657084, 264760, 264448, 264288, 264767, 264689, 21906766, 21906767, 21906767, 21906769, 255021, 265022, 60170615, 33657023, 27486265, 35696423, 35696423, 224639, 87168518, 22279002, 264563, 264886, 18108394	264259, 29331826, 264508, 264908, 264510, 264007, 285011, 264288, 264637, 18108385	264592	29331825, 265017, 265018, 264288, 265020, 265021, 264634, 56526486	264768, 22278997, 265021, 264690, 264259, 264692, 29331824, 26331828, 264509, 264907, 29331828, 264509, 264909, 265007, 265009, 264628, 20281069, 264909, 265007, 265009, 264632, 264636, 264591, 264592, 264639, 265018, 265019, 22779002, 26463, 264648, 264448, 264684, 264687, 264685, 264684, 264448, 264684, 264687, 264685, 264684, 264448, 264684, 264687, 264685, 264684, 264448, 264684, 264687, 264	56182575, 264092, 29331824, 29331826, 29331830, 265017, 265018, 265020, 83373044	22278996, 22278997, 22278999, 60432049, 29331822, 29331824, 33657402, 85658542, 265011, 265018, 265019, 21906767, 21906768, 264693, 18108385, 22279000, 22279002	264907, 265019	56182575, 22278996, 22278997, 264259, 29331822, 29331825, 264509, 264112, 265009, 264593, 21906754, 265018, 265019, 264448, 264288, 264685, 21906769, 265020, 265022, 264691, 18108370, 65274791, 264553	265018, 264763, 264683, 264691
	₹ê	UNCLASSIFIED	ubiquitin	UNCLASSIFIED	UNCLASSIFIED	eph	collagen	transport	UNCLASSIFIED
			Contains protein domain (PF00628) - ubiquitin PHD-finger	Contains protein domain (PF00098) - UNCLASSIFIED Zinc finger, CCHC dass	Contains protein domain (PF01363) - UNCLASSIFIED FYVE zinc finger	Contains protein domain (PF00431) - eph CUB domain		Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	
Novel Protein sim. GBank gi[5138930]gb AAD40382.1 - (AF093680) transcription factor IIB [Homo sapiens]			87539364 (4155, 4156) Novel Protein sim. GBank gil4220590 dbj BAA74579 - (D87908) nuclear protein np95 [Mus musculus]		94136689 (4159, 4160) Novel Protein sim. GBank gi 2408021 emb CAB16219.1 - (Z99162) putative vacuolar protein [Schizosaccharomyces [pombe]	94847186 (4161, 4162) Novel Protein sim. GBank gil5524734[gb AAD44360.1 AF16635 - (AF166350) ST7 protein [Homo sapiens]	87628629 (4163, 4164) Novel Protein sim. GBank gij3880558jemb CAA94234j - (Z70271) predicted using Genefinder; similar to collagen; CDNA EST yk308e7.3 comes from this gene; cDNA EST yk308e7.5 comes from this gene; cDNA EST yk385a8.3 comes from this gene; cDNA EST yk385a8.3 this gene [Caeno	94141000 (4165, 4166) Novel Protein sim. GBank gi 2352427 (AF004161) - peroxisomal Ca-dependent solute carrier [Oryctolagus cuniculus]	95199298 (4167, 4168) Novel Protein sim. GBank 911728336 sp P39193 ALU6_HUMAN - III! ALU SUBFAMILY SP WARNING ENTRY III!
	5 87594118 (4151, 4152)				94136689 (4159, 4160)	94847186 (4161, 4162)	87628629 (4163, 4164)		95199298 (4167, 4168)
207	2076	2077	2078	2079	2080	2081	2082	2083	2084

2085	94989476 (4169, 4170)	2085 94989476 (4169, 4170) Novel Protein sim. GBank gi 1655699 emb CAA69032 -	UNCLASSIFIED	56182575, 60432289, 264908, 56182435
		(TU//52) pherophorn-5 [Volvox carlen]		87168474, 264763, 264369, 264688, 264693, 18108370, 56182323
2086	91234404 (4171, 4172)	91234404 (4171, 4172) Novel Protein sim. GBank gij3875032jemb CAA88936j -	UNCLASSIFIED	35696286, 264259, 35696052, 264906.
		(249125) Similanty to Inchostrongylus colubriformis 11 kd		264907, 264908, 264909, 264910, 264759,
		Secretary protein (SWISS First accession number P21937);		264604, 264762, 264768, 264769, 35695917,
		FST FMRI - D37844 comes from this good: DNA FST		263978, 35696423, 35695855, 264632,
		EMBL: D36149 come		264634, 264637, 264638, 264639, 56182323,
2087	21436337 (4173, 4174)	-		16106363, 264482, 264486
,	84111527 (4175 4176)	94111527 (4175, 4176) Novel Protein cim CBank Allasanooniams (CAA46234.41	UNCLASSIFIED	264489
	(a	(Al 021481) similar to Described incomparations and		264488, 22278994, 35696286, 22278996,
		Check the second		29331827, 35696052, 33657402, 21906754,
		EMBI : 026469 compo form this company form		33109954, 87168474, 265017, 265018,
		EMBL: Date Comes from this gene; CDIVA ES	_	285019, 264448, 264683, 264369, 264685.
_		EmbL.D/069/ comes from this gene; cDNA EST yk373h9.5	-	264687, 264689, 21906765, 21906766,
	•	comes from this gene; cDNA EST EMBL: 10080		21906767, 21906768, 21906769, 265020,
				265021, 265022, 264692, 33657023,
				33657109, 33657182, 27486261, 27486262,
				33657349, 27486265, 35696423, 35695855,
0000	05400004 (4477 4470)			83373044, 87168518, 22279000, 264567
	93422001 (4177, 4178)		cadherin	18108392, 264488, 52644507, 18108394,
		gila/35116jreflNP_004623.1pDAP3 - Death associated		18108397, 52646842, 18108398, 56182575,
		protein 3		22278994, 22278995, 35696286, 22278996,
				56994075, 22278997, 22278998, 22278999,
				264091, 264092, 264093, 264094, 60432049,
			•	264259, 29331822, 20281099, 29331824,
				29331825, 29331826, 29331827, 29331828,
				35696052, 33656970, 29146498, 29146499.
			_	264102, 264106, 264107, 264109, 264508,
				264905, 264509, 264906, 264907, 264908,
				66712502, 264828, 52644045, 264909,
				56182435, 264110, 264112, 264510, 264511,
				265006, 264512, 265007, 265008, 264910,
_				265009, 60170831, 264592, 264593,
				60433356, 33657402, 60433438, 264595,
				55812038, 264758, 21906754, 33657084,
				55811386, 52644296, 265010, 265011,
_				87168559, 265017, 265018, 265019, 264760,
				264761, 55811150, 264762, 18108351,
				264682, 264448, 264763, 264764, 264683,
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				18108358, 56181562, 264769, 18108359,
		-		264689, 21906765, 21906766, 21906767,
				21906768, 29148627, 21906769, 55811957,
				29148629, 29148784, 35695917, 265020.
				265021, 265022, 60170615, 264690,

2090	2090 88222470 (4179, 4180)				22278006 22278008 22278000 264360
					29331826, 35696052, 264910, 33657402,
					60433438, 33109954, 87168474, 87168559,
					265018, 265019, 264681, 264684, 264686,
					264687, 264688, 264689, 21906765,
					21906766, 21906767, 21906769, 35695917,
					265022, 60170615, 33657023, 35696423,
,	06200164 /4484 44821				35695855, 264952, 18108387, 22279000
E0.7	(90309101 (4181, 4182)			UNCLASSIFIED	263994, 264905, 264908, 264511, 264512,
		gif4580997[gb]AAD24571.1[AF12108 - (AF121081) CAMP			265008, 264910, 55811386, 264288, 264768,
		inducible 2 protein [Mus musculus]			56181562, 21906765, 21906768, 21906769,
					265022, 264628, 264563, 264567
2082	88223605 (4183, 4184)			ротеорох	22278997, 22278999, 66712502, 87168559,
					264683, 265021, 264486
2093	87406073 (4185, 4186)	87406073 (4185, 4186) Novel Protein sim. GBank gi 2352427 (AF004161) -	Contains protein domain (PF00153) - transport	transport	264905, 264906, 264907, 264908, 264510,
		peroxisomal Ca-dependent solute carrier [Oryctolagus	Mitochondrial carrier proteins		265006, 265007, 265009, 264910, 264596,
		cuniculus)			21906754, 87168474, 265011, 264603,
					265018, 265019, 264760, 264766, 264768,
					264769, 21906767, 21906768, 21906769,
					265021, 264690, 33657023, 264693, 264628.
					264634, 264636, 264637, 264557, 56182323,
					264564
2094	(91230929 (4187, 4188)	91230929 (4187, 4188) Novel Protein sim. GBank		MHC	35696286, 265017, 265018, 265019
		gil4929551 gb AAD34036.1 AF15179 - (AF151799) CGI-40			18108388
1000	_	protein (Homo sapiens)			
2082		95351526 (4189, 4190) Novel Protein sim. GBank gi 1363238 pir A57284 -	Contains protein domain (PF00035) - dna_ma_bind	dna_ma_bind	35696286, 52644045, 265006, 265007,
		spermatid perinuclear RNA-binding protein Spnr - mouse	Double-stranded RNA binding motif		265008, 87168559, 18108351, 21906769.
					29148784, 265020, 33657023, 27486262,
					18108374, 18108388
2096	94119760 (4191, 4192)	94119760 (4191, 4192) Novel Protein sim. GBank gi 3834423 (AF070689) -	Contains protein domain (PF00400) -	ATPase_associated	264488, 264489, 65274572, 56182575,
		cytoplasmic dynein intermediate chain isoform DIC1a	WD domain, G-beta repeat	1	22278996, 22278997, 22278999, 264259,
		[Drosophila melanogaster]			60432289, 29331826, 35696052, 264107,
					264508, 264509, 264905, 264906, 264907,
					264908, 52644045, 264909, 264510, 264511,
					264512, 265008, 264910, 265009, 264592,
					60433356, 60433438, 264758, 264596,
					55812038, 21906754, 264601, 264602,
					264605, 264762, 264681, 18108351, 264764,
					264683, 264288, 264687, 264768, 264769,
					264689, 21906765, 21906766, 21906767,
					35695917, 265020, 265022, 52644150,
					264691, 264692, 33657023, 264693,
					27486261, 35695763, 264628, 264629,
					35696423, 35695855, 264631, 264632,
					264634, 264635, 264555, 264637, 263981,
					264638, 264639, 264563, 264483, 264565,
					264566, 264486, 264567

2097	95322772 (4193, 4194)	2097 95322772 (4193, 4194) Novel Protein sim. GBank	Contains protein domain (PF00096) - transcriptfactor	transcriptfactor	65274572 264511 265010 264600 265017
		gi 5174501 ref NP_006051.1 pLYF1 - zinc finger protein, subfamily 1A, 1 (lkaros)	Zinc finger, C2H2 type		264448, 264288, 265021, 60170615, 264692,
2098		87780340 (4195, 4196) Novel Protein sim. GBank	Contains protein domain (PE00782)	phoenhalaeo	5303/109, 181083/0, 264636, 264483
		gil4758208freflNP_004081.1 pDUSP - dual specificity	Dual specificity phosphatase.	acena de cond	30334073, 204239, 204288, 203020, 204363
2000		osariosoz (4107 4108) Nicel Barrier (CB)	catalytic domaín		
		prover Frotein sim. Gbank gijzb95659 (AF026954) - pyruvate dehydrogenase phosphatase regulatory subunit		phosphatase	65274572, 264905, 65274444, 264691, 264636, 264555
2100		pegages (4100, 4200) Mecursor; PDPr [Bos taurus]			
3		Novel Protein sim. GBank gij3881189jemb CAB16514 - 799281 similar to ADP-ribosofation factor, cONA ECT	Contains protein domain (PF00025) - nucl_recpt	nucl_recpt	56182575, 22278995, 22278996, 22278997,
_		[EMBI :CO8179 comes from this need: conv. cc.t.	AUP-ribosylation factor family		22278998, 60432049, 264259, 29331822,
		EMBL: C08337 comes from this gene: cDNA EST			29331824, 29331825, 29331827, 29331828,
		EMBL: C09829 comes from this gene; cDNA EST vk291b4.5	100		29146498, 264909, 265008, 265009, 264910, 1264604
		comes from this gene; cDNA EST vk4			204331, 00432229, 60433336, 3365/402,
					204738, 2190b/34, 83638342, 87168474,
					265017, 265018, 265019, 264681, 18108351,
					264762, 264448, 264369, 264288, 18108355,
					264686, 21906765, 21906767, 21906768.
					21906769, 265020, 265021, 33657023,
					18108374, 35696423, 264558, 83373044,
2101	R7782604 (4201 4202)	Maria Destate all Control of the Con			87168518, 60432113, 22279000, 22279002
	(1076) (1701) 1702)	(120) (120), 120) (120)		UNCLASSIFIED	264091, 29331824, 264105, 265007, 265010.
2102	87770461 (4203, 4204)	2102 87770461 (4203, 4204) Novel Protein sim. GBank oil 3874149 emhi CAA97423 11.			100300
		(273103) predicted using Genefoder (Caeparte Aditio		UNCLASSIFIED	264488, 264489, 35696286, 264259,
		(2.5.5.5) predicted using denember (Caenornapolitis			35696052, 264508, 264905, 264907, 264908,
					264909, 264511, 264512, 264591, 264593,
					60433356, 264758, 264601, 264605, 264760,
					18108351, 264448, 264764, 264288, 264767,
					264768, 21906769, 35695917, 18108374,
					264634, 264555, 264559, 264563, 264482,
2103	95413576 (4205 4206)	95413576 (4205 4206) Novel Protein sim CBank alt 210310150141:184471059 11			264486
	(2011)	- IL 9097874838.11	Contains protein domain (PF01530) - transcriptfactor	transcriptfactor	65274572, 56994075, 22278999, 264259,
		(Activities of the same of the	Zinc finger, C2HC type		29331824, 29331825, 35696052, 29331828.
			_		66712502, 265009, 60170831, 264595,
					33109954, 85658542, 87168559, 265017,
					265019, 264448, 21906765, 21906768,
					265022, 33657023, 27486262, 33657349,
					35695763, 60431528, 18108374, 55811576,
					56182323, 18108387, 87168518, 60432113,
2104	2104 85776161 (4207, 4208)				264564
	7			UNCLASSIFIED	264592, 264604, 22279000

264488, 56182575, 22278994, 56994075, 22278996, 22278996, 22278999, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 20331824, 29331826, 60432289, 29331827, 29331826, 294598, 264598, 264599, 264599, 264599, 264599, 264599, 264599, 264599, 264599, 264599, 264762, 18108351, 265017, 265018, 265019, 264762, 18108351, 264684, 18108354, 264288, 264686, 21906766, 21906767, 21906768, 21906769, 21906769, 265022, 60170815, 52644150, 265020, 265022, 60170815, 52644150, 264692, 33657029, 264693, 33657109, 60431528, 18108374, 66274791, 35695855, 264635, 60170334, 264569, 264568, 81108385, 18108385, 264568, 81168518, 60432113, 264566, 264566, 81168518, 60432113, 264566, 264566, 81168518, 60432113, 264566, 264566, 81168518, 60432113, 264566, 264566, 81168518, 60432113, 264566, 264566, 81168518,	265006, 265019	264906, 264639	264905, 264906, 264907, 264908, 264909, 264758, 265011, 264600, 264601, 264764, 264766, 264769, 26469, 264693, 264639, 3569585, 264632, 264634, 264635, 264638, 264486	18108348, 264769, 18108370, 18108374, 264555, 264556, 264557, 264558	264564	264766, 35695917, 264630, 264567, 264486	264508, 264906, 264591, 264682, 22279002	265006	66714117, 29331826, 29331827, 60433438, 55812038, 265017, 265019, 264689, 21906769, 55811957, 265020, 265021, 33657109, 60170394, 264558	65274572, 264689, 264691, 264692, 60432113
UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	struct	cadherin		UNCLASSIFIED
							Contains protein domain (PF00069) - struct Eukaryotic protein kinase domain	Contains protein domain (PF00028) - cadherin Cadherin		
2105 94848080 (4209, 4210) Novel Protein sim. GBank gij1707032 (U80445) - coded for by C. elegans CDNA yk13g5.3; coded for by C. elegans CDNA yk21g6.3; coded for by C. elegans CDNA CEMSE18F; coded for by C. elegans cDNA yk126b1.3; coded for by C. elegans CDNA yk65h8.3; coded for by C. elegans CDNA yk65h8.3; coded for by C. elegans CDNA yk65h8.3.	4212)	79822662 (4213, 4214) Novel Protein sim. GBank gi 3881524 emb CAA93883 - (270038) ZK1067.4 [Caenorhabditis elegans]	94233976 (4215, 4216) Novel Protein sim. GBank gil3176689 (AC003671) - Contains similarity to ubiquitin carboxyl-terminal hydrolase 14 gb Z35927 from S. cerevislae. [Arabidopsis thaliana]	4218)	87729075 (4219, 4220) Novel Protein sim. GBank gil481043 pir S37671 - bat2 protein - human	4222)	87293783 (4223, 4224) Novel Protein sim. GBank gi 2143639 pir 156542 - calmodulin-binding protein - rat	78941388 (4225, 4226) Novel Protein sim. GBank gil4426629 gb AAD20459 - (AF100960) protocadherin [Rattus norvegicus]	87889342 (4227, 4228) Novel Protein sim. GBank gij3327184 dbj BAA31660 - (AB014585) KIAA0685 protein [Homo sapiens]	2115 90993785 (4229, 4230) Novel Protein sim. GBank gil4757890 refiNP_004328.1 pC8OR - chromosome 8 open reading frame 1
94848080 (4209, 4						87818419 (4221, 4222)				90993785 (4229, 4
2105	2106	2107	2108	2109	2110	2111	2112	2113	2114	2115

4534)	2116 86259387 (4231, 4232) Novel Protein sim. GBank gi 2246532 (U93872) - ORF 73,		struct	35696286, 22278999, 56182181, 29331824,
assoc	associated herpesvirus]			29331023, 29331027, 33098032, 264907, 56182435, 265008, 264591, 55812038,
				55811386, 87168559, 264288, 264369, 21906769, 29148629, 33657023, 35695763
				55811576, 35696423, 18108385
Š	87788904 (4233, 4234) Novel Protein sim. GBank gi 2330021 (AF019250) - kinesin- Iralated protein: VPB- Contact (Proceeding Technology)		struct	29331824, 264511, 265009, 33109954,
	co protein, and , costait [crosopinia metahogaster]			253017, 253018, 264288, 264689, 253020, 264692, 56526486, 264482
ĝ.	87078894 (4235, 4236) Novel Protein sim. GBank gij1079307[pir][B56573 - nuclear		glycoprotein	264259, 264905, 264907, 264908, 264510,
	pore compiex glycoprotein p62 - African clawed frog			264511, 265009, 264910, 265010, 264602, 264288, 264768, 264693, 263967, 263972,
S S	86999317 (4237, 4238) Novel Protein sim. GBank gil4321407[gb]AAD15748 - (AF047690) ATP-binding cassette protein M-ABC1 [Homo		UNCLASSIFIED	264693, 18108385
	87789395 (4239, 4240) Novel Protein sim CBank	(C. 2000) C.		
9.48	Nover Frotein suit. Oberink gij4885527/refINP_005480.1 pNSP3 - novel SH2-containing Src homology domain 2	Contains protein domain (PF00017) - epn Src homology domain 2	ebu	264091, 264259, 29331826, 29331828. 265017, 264604, 264288, 264685, 265020.
prot				264691, 18108370, 55810764, 264555, 264636, 60432113
Nov 19147	80021375 (4241, 4242) Novel Protein sim. GBank gil4757728 ref NP_004886.1 pAGTA - andiotensin/vasopressin recentor All/AVP-like		UNCLASSIFIED	264601, 264766, 263978
ĺĝ	91230931 (4243, 4244) Novel Protein sim. GBank			18108394, 56182575, 22278997, 29331822,
9 <u>i</u>	gil4929551[gb AAD34036.1]AF15179 - (AF151799) CGI-40			29331824, 29331825, 29331826, 29331828,
D D	protein [Homo sapiens]			264907, 56182435, 265007, 264910, 265010,
				203016, 204000, 203020, 33011370, 204333. 264637, 18108382, 83373044, 18108383.
				18108384, 56526486, 264565, 264567
Nov	86787998 (4245, 4246) Novel Protein sim. GBank gi 2224551 db BAA20764 -	Contains protein domain (PF01363) - struct	struct	18108396, 264757, 265011, 18108351,
킨:	(ABUUZ3U3) KIAAU3U3 (Homo sapiens)	FYVE zinc finger		264691, 264634, 18108385
é 8	83005951 (4247, 4248) Novel Protein sim. GBank gi 5689455 db BAA83011.1 - (AB028982) KIAA1059 protein [Homo sapiens]	Contains protein domain (PF00801) - transport PKD domain	transport	29331822, 264906, 264907, 264591, 264639, 264563
£	95354041 (4249, 4250) Novel Protein sim. GBank		UNCLASSIFIED	264259, 264509, 264907, 264511, 85658542,
5	gij728831[sp P39188 ALU1_HUMAN - !!!! ALU SUBFAMILY J WARNING ENTRY !!!			264763, 21906765, 35695917, 264636, 264488
12	95084231 (4251, 4252) Novel Protein sim. GBank gil4539264 emb CAB39853.1 -		UNCLASSIFIED	264488, 264489, 29331827, 35696052,
₹	(AL049495) conserved hypothetical protein			264905, 264509, 264908, 264909, 264510,
တ္တ	[Schizosaccharomyces pombe]			265009, 264591, 264592, 264593, 33657402,
				264594, 264595, 264596, 264758, 264601,
				264603, 265018, 264604, 264605, 264760,
				264681, 264762, 264583, 264764, 264684,
				264288, 264685, 264689, 60170615,
				5505/025, 5505/109, 55610/64, 204055, 264636 264637 264638 264639 83373044
				264564, 264566

35696286, 29331826, 35696052, 264508, 264509, 264905, 264906, 264907, 264909, 264909, 264907, 264908, 264909, 264907, 264908, 265007, 265009, 264910, 264768, 265011, 264600, 264601, 264604, 264769, 264689, 264569, 264691, 264691, 264691, 264691, 264692, 264693, 264631, 264631, 264631, 264631, 264631, 264631, 264631, 264639, 18108374, 3569565, 264631, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264567	56182575, 35696286, 56182181, 29331824, 60432289, 35696052, 264905, 264907, 66712502, 264908, 264909, 264510, 265009, 264910, 264591, 55812038, 265018, 264764, 264288, 264369, 264687, 264768, 55811957, 264692, 18108368, 264628, 264634, 264635, 264637, 56182323, 264639, 18108388, 264563, 264563, 264563	66714117, 264828, 264595, 55812038, 55811150, 55811957, 264693, 18108374, 263978, 65274791, 18108381, 83373044, 22279000	263981 35696052, 264909, 264768, 35695917	Contains protein domain (PF00122) - ATPase_associated 264488, 22278999, 264559, 29331827, 29331828, 35696052, 264509, 264905. 264906, 264907, 264908, 264909, 264905, 265006, 264511, 265007, 26490, 264910, 265500, 26451, 265007, 264910, 264600, 264511, 265007, 264910, 264600, 264601, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264600, 264629, 264769, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264639, 264555, 264631, 264634, 264565, 264630, 264565, 264566, 264565, 264566, 264565, 264566, 264565, 264566, 264565, 264566, 264565, 264566, 264567
	A	UNCLASSIFIED	UNCLASSIFIED polassium_channel	ATPase_associated
			UNCLASSIFIED Contains protein domain (PF00805) - potassium_channel Pentapeptide repeats (8 copies)	Contains protein domain (PF00122) -
2127 81118652 (4253, 4254) Novel Protein sim. GBank gi 4868435 gb AAD31315.1 AF14323 - (AF143236) apoptosis related protein APR-2 [Homo sapiens]			9	95361096 (4263, 4264) Novel Protein sim. GBank gil5689373 dbj BAA82973.1 - , (AB028944) KIAA1021 protein [Homo sapiens]
81118652 (4253, 4254)	87414262 (4255, 4256)	95102089 (4257, 4258)		
2127	2128	2129	2130	2132

60424179, 52646365, 52646842, 56994075, 35696286, 22278998, 66432049, 56182181, 66714117, 60424269, 29331826, 29331828, 35696052, 264905, 264906, 264907, 66712502, 29331830, 56182435, 265006, 264512, 265008, 60431735, 60433356, 33657402, 55812038, 33109954, 21906754, 55811386, 255010, 264603, 265017, 265018, 265010, 264603, 265017, 265018, 265019, 55811150, 18108351, 265020, 265021, 60170615, 356595917, 265020, 265021, 60170615, 3565923, 33657109, 60431528, 18108374, 35695423, 6518380, 56182323, 83373044, 18108385, 18108387, 60432113, 222799002, 264563, 264566	56181686, 35696286, 21906754, 55811386, 265011, 265017, 18108351, 264765, 264766, 264688, 21906768, 35695917, 265020, 33657023, 264628, 35695855, 264632, 264555, 264556, 264557, 264558, 18108382, 22279002	22278999, 29331828, 35696052, 264906, 264908, 264908, 2649010, 265009, 264591, 264758, 52646317, 265011, 87168559, 264601, 18108351, 26448, 264683, 264684, 264689, 18108359, 264691, 33657023, 264692, 35695763, 264636, 264637, 56182323, 264639, 22279002, 264564	264639	264569, 264909, 33109954, 264763, 21906768, 60170394, 18108385, 264563		264905, 264910, 264591, 55812038, 55811386, 85658542, 264760, 18108351, 18108359, 55811957, 265020, 265021, 33657023, 18108364, 55811576, 83373044, 18108385, 56526486, 264482
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	ATPase_associated	
		Contains protein domain (PF00628) - UNCLASSIFIED PHD-finger				
2133 95351539 (4265, 4266) Novel Protein sim. GBank gil4220489 (AC006069) - hypothetical protein [Arabidopsis thaliana]	95412697 (4267, 4268) Novel Protein sim. GBank gij3875351 emb CAB09415 - (296047) DY3.6 [Caenorhabditis elegans]	88079813 (4269, 4270) Novel Protein sim. GBank gil5689559 dbj BAA83063.1 - (AB029034) KIAA1111 protein [Homo sapiens]	84346479 (4271, 4272) Novel Protein sim. GBank gil2662167 dbj BAA23715 - (AB007903) KIAA0443 [Homo sapiens]	87637716 (4273, 4274) Novel Protein sim. GBank gil4884110 emb CAB43262.1 - (AL050090) hypothetical protein [Homo sapiens]	87395446 (4275, 4276) Novel Protein sim. GBank gil5174779 gb AAD40696.1 - (U87804) 50 kDa protein [Caulobacter crescentus]	94843882 (4277, 4278) Novel Protein sim. GBank gij3850821 jemb CAA77135j - (Y18350) U2 snRNP auxiliary factor, large subunit [Nicotiana plumbaginifolia]
133 95351539 (4265, 4266)		2135 88079813 (4269, 4270)		2137 87637716 (4273, 4274)		2139 94843882 (4277, 4278)

264488, 264259, 29331824, 264104, 264109, 264509, 265006, 264759, 265018, 264448, 264288, 21906768, 55811957, 265021, 33657023, 27486265, 35696423, 264636, 264556, 264559, 264566	265020, 264693	263978	22278997, 29331827, 264907, 265020, 60432113	65274572, 264259, 29331824, 29331827, 264906, 264508, 264591, 87168559	264600, 265019, 284288, 264768, 21906765, 21906765,	22279002	263978	264909, 60433356, 264686	264907, 264768, 264769, 18108385		264593	29331824 29331826 35696052 264758	87168474, 265018, 52644150, 33657109	22278998, 29331822, 29331824, 29331828, 264764, 264769, 21906766, 264486	22278996, 56994075, 22278999, 60432049	264259, 29331822, 29331824, 29331826,	35696052, 29331828, 264508, 264511,	60174639, 265010, 265011, 87168559,	265017, 265018, 265019, 264448, 264288,	264689, 21906765, 21906766, 21906768.	265020, 60170615, 33657109, 33657182.	33657349, 18108370, 264635, 264557,	1001/0394, 10100303, 0/ 100310, 222/9000	16106397, 16106396, 263007, 264391, 265011, 18108351, 18108368, 18108374, 18108388
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	alycoprotein		UNCLASSIFIED				_			~		o de dia	as per la
												Contains protein domain (PF00059) - alycoprotein	Lectin C-type domain											
2140 87645655 (4279, 4280) Novel Protein sim. GBank gil4417293 gb AAD20418 - (AC007019) unknown protein (Arabidopsis thaliana			94140051 (4285, 4286) Novel Protein sim. GBank gi 2135766 pir S53362 - mucin 5AC (done JER47) - human (fragment)	94320114 (4287, 4288) Novel Protein sim. GBank gi 2078483 (U43200) - antifreeze glycopeptide AFGP polyprotein precursor Boreogadus	saida)			87010515 (4291, 4292) Novel Protein sim. GBank gi 1255871 (U53341) - short region of weak similarity to bovine membrane receptor p63	80432911 (4293, 4294) Novel Protein sim. GBank gij3080398jembjCAA18718.11 -	(AL022603) putative protein [Arabidopsis thaliana]	80048811 (4295, 4296) Novel Protein sim. GBank gi 728837 sp P39194 ALU7_HUMAN - IIII ALU SUBFAMILY SQ WARNING ENTRY IIII	Novel Protein sim. GBank	gil 19863 sp P20693 FCE2_MOUSE - LOW AFFINITY IMMUNOGLOBULIN EPSILON FC RECEPTOR (LYMPHOCYTE IGE RECEPTOR) (FC-EPSILON-RII) (CD23)	94140059 (4299, 4300) Novel Protein sim. GBank gij5420387jembjCAB46579.1j - (AJ243459) proteophosphoglycan [Leishmania major]	95353241 (4301, 4302) Novel Protein sim. GBank gi 5689407 dbj BAA82987.1 -	(AB028958) KIAA1035 protein [Homo sapiens]							79321640 (4303 4304) Novel Protein sim GBank ni12452473 (AED84205)	serine/threonine protein kinase TAO1 [Rattus novegicus]
87645655 (4279, 4280)	79623986 (4281, 4282)	80041222 (4283, 4284)					20564305 (4289, 4290)	87010515 (4291, 4292)	80432911 (4293, 4294)		80048811 (4295, 4296)	87362022 (4297, 4298)		84140059 (4299, 4300)	95353241 (4301, 4302)				-				79321640 (4303 4304)	
2140				2144			2145	2148	2147]	2148	2149			2151								2152	

2153	88313371 (4305, 4306)	2153 88313371 (4305, 4306) Novel Protein sim. GBank gil4758704 reflNP_004216.1 pMASL - MFH-amplified sequences with leucine-rich tandem repeats 1	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat		264488, 263994, 52646842, 22278996, 22278998, 22278999, 264259, 29331822, 35696052, 264508, 264509, 264905, 264906, 264906, 264907, 264907, 264908, 264909, 5618243, 264510, 264511, 264512, 264758, 87188474, 87168559, 265017, 265019, 264760, 264288, 2040566, 21906768, 35695917, 33657023, 33657109, 35695855, 264631, 264632, 264634, 264488, 264566, 264488
2154	ı	87408034 (4307, 4308) Novel Protein sim. GBank gi 225150 pri 1209265U - chorion protein B11 [Bombyx mori]		UNCLASSIFIED	56994075, 264094, 265009, 265019, 264288, 21906767, 35695917
2155	87424072 (4309, 4310)			UNCLASSIFIED	18108392, 18108398, 22278996, 264259, 29331824, 265008, 265010, 265011, 265017, 265019, 264689, 264686, 265020, 264693, 264628, 56182323
2156		84295205 (4311, 4312) Novel Protein sim. GBank gil3970966 (AC004974) - spa-1- like; similar to AF026504 (PID:92555183) [Homo sapiens]			265007, 264684
					264591
		86444218 (4315, 4316) Novel Protein sim. GBank gij1076211 pir S50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardtii		Ω	264596
		80083729 (4317, 4318) Novel Protein sim. GBank gil4650844 dbj BAA77027.1 - (AB026190) Kelch motif containing protein [Homo sapiens]	Contains protein domain (PF00651) - dna_rna_bind BTB/PO2 domain		29331822, 264112, 265009, 264691, 33657023, 264634
					264634
2161				UNCLASSIFIED	265008
2162		94319526 (4323, 4324) Novel Protein sim. GBank gil1504006 dbj BAA13202 - (D86966) similarto human ZFY protein. [Homo sapiens]			65274572, 264508, 264905, 264906, 264907, 264908, 5264045, 264909, 265007, 264910, 264591, 264591, 264591, 264591, 264591, 264591, 264600, 264762, 264763, 264683, 264764, 264689, 265020, 264691, 264628, 264699, 264628, 264691, 264628, 264699, 264659, 264691, 264658, 264639, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 264659, 26456, 26456, 264569, 26456
26	854 (1) 50 (4,325, 4,326)	934 I 7 130 (4323, 4340) Novel Protein sim. GBank gij38 (6537 jembjCAA98270) - (273974) cDNA EST yk291f5.3 comes from this gene; cDNA EST yk291f5.5 comes from this gene [Caenorhabditis elegans]		SIFIED	56182575, 22278996, 264093, 264683, 33657023, 65274620, 60432113
2164	80569456 (4327, 4328)		Contains protein domain (PF01006) - collagen Hepatilis C virus non-structural protein NS4a		264603, 264637, 264565

56994075, 22278996, 22278997, 22278999, 264259, 29331822, 29331824, 29331825, 29331826, 29331826, 29331826, 29331827, 29331828, 264906, 29331830, 56182435, 265009, 21906754, 33657084, 265011, 265019, 264448, 264288, 265360, 21906769, 265360, 21906769, 26536, 21906769, 26566, 21906769, 219	52645156, 22278994, 22278998, 65714117, 29331828, 52544045, 265018, 265019, 264369, 21906765, 21906767, 21906768, 21906769, 265021, 265022, 264693, 27486262, 25695763, 18108376, 55526486, 87168518, 264567	56182575, 35696286, 29331824, 29331826, 29146498, 56182435, 265008, 265009, 264592, 264592, 33657402, 33109954, 265011, 265017, 265018, 18108351, 264369, 21906764, 21906765, 21906768, 29148627, 21906769, 52644150, 33657109, 35696423, 18108381, 18108384, 18108385, 60432113, 264557	66714117, 29331827, 264907, 264511, 264591, 265018, 264764, 264683, 264766, 264768, 264566	264629, 264555, 264559	65274572, 56182575, 22278997, 22278998, 264259, 29331825, 264509, 264906, 56182435, 60433438, 55812038, 264596, 55811386, 265019, 264762, 264763, 26448, 264764, 264684, 264762, 264766, 264685, 56181562, 264689, 55811957, 265200, 264535, 264689, 55811957, 265200, 264534, 264689, 55811576, 66431528, 18108374, 35696423, 55811576, 65274791, 264634, 264659, 264558, 87168518, 60432113, 264564	264369, 265020, 264558	264259, 264558	264369	264906, 35695855, 264555, 264557
UNCLASSIFIED	ubiquilin	UNCLASSIFIED	transport		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	
·			Contains protein domain (PF00664) - transport ABC transporter transmembrane region.						
2165 94329169 (4329, 4330) Novel Protein sim. GBank gi 1086794 (U41107) - Nodefinition line found [Caenorhabditis elegans]	87618934 (4331, 4332) Novet Protein sim. GBank gi 2706522 emb CAA75816 - (Y15895) ubiquitin activating enzyme [Drosophila melanogaster]	87716864 (4333, 4334) Novel Protein sim. GBank gi[2224713 dbj BAA20840] - (AB002384) KIAA0386 [Homo sapiens]	86999334 (4335, 4336) Novel Protein sim. GBank gil4321407 gb AAD15748 - (AF047690) ATP-binding cassette protein M-ABC1 [Homo sapiens]		94141033 (4339, 4340) Novel Protein sim. GBank gil5106521[gb]AAD39741.1[AF10536 - (AF105365) K-Cl cotransporter KCC4 [Homo sapiens]		85452460 (4343, 4344)	Novel Protein sim. GBank gil4309681 gb AAD15478 - (AC006930) R33423_1 [Homo sapiens]	95003288 (4347, 4348) Novel Pratein sim. GBank gi 2493778 sp Q09456 YQ35_CAEEL - PUTATIVE CUTICLE COLLAGEN C09G5.5
5 94329169 (4329, 4330 <u>)</u>	2166 87618934 (4331, 4332)	2167 87716864 (4333, 4334)	2168 86999334 (4335, 4336)	2169 87886937 (4337, 4338)	94141033 (4339, 4340)	2171 80194050 (4341, 4342)			2174 95003288 (4347, 4348)

2175	94325850 (4349, 4350)	2175 94325850 (4349, 4350) Novel Protein sim. GBank gij1263287 (U47855) - fibroin-3 [Araneus diadematus]		UNCLASSIFIED	264488, 35696286, 20281099, 29331826, 60432289, 35696052, 264109, 264508, 264509, 264907, 264908, 2649
					264909, 264510, 264511, 265006, 265007, 264512, 265008, 264910, 264591, 264594, 264595, 264596, 264758, 55812038, 265011,
					264600, 264603, 264760, 264762, 264448,
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					264628, 264629, 55811576, 35696423,
					65274791, 35695855, 264631, 264632,
					264634, 264635, 264636, 264637, 264638,
					264639, 264558, 18108385, 60432113,
					264563, 264564, 264565, 264566, 264486,
_					264567
2176	88223392 (4351, 4352)	88223392 (4351, 4352) Novel Protein sim. GBank	Contains protein domain (PF00805) - oncogene	oncogene	52644507, 52646842, 22278994, 35696286,
	•	_	HUMAN - !!!! ALU SUBFAMILY Pentapeptide repeats (8 copies)	1	22278996, 22278999, 29331826, 29331827,
		SQ WARNING ENTRY !!!			35696052, 29331828, 33656970, 29331830,
					264910, 33657402, 264758, 52644296,
					87168559, 265018, 264689, 21906765.
					21906767, 21906769, 35695917, 52644150,
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					33657182, 27486261, 27486262, 33657349,
					18108376, 18108377, 35695855, 87168518,
					60432113, 264404, 22279000, 264486
2177	94128942 (4353, 4354)	94128942 (4353, 4354) Novel Protein sim. GBank		kinase	18108392, 22278997, 22278999, 264093,
		gi 5454072 ref NP_006416.1 pSLU7 - step II splicing factor			33657402, 265019, 264448, 264766, 264689,
_		SLU7			21906767, 21906768, 21906769, 265021,
					33657023, 18108370, 18108374, 60432113,
					22279002
2178	_	87601557 (4355, 4356) Novel Protein sim. GBank gil473407 (U08215) - NST-1 [Mus	gil473407 (U08215) - NST-1 [Mus Contains protein domain (PF00012) - eph	ude	264488, 22278996, 22278999, 29331824,
		musculus]	Hsp70 protein		29331825, 29331826, 29331827, 52644296,
					87168474, 18108370, 35695855, 22279002
2179	2179 87316275 (4357, 4358)			UNCLASSIFIED	60424269, 264760, 264628, 264632

2180	95351397 (4359, 4360)	2180 95351397 (4359, 4360) Novel Protein sim. GBank gij3122317jspjP90648jKMHB_DiCDI - MYOSIN HEAVY CHAIN KINASE B (MHCK B)	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		52644507, 22278994, 35696286, 22278997, 22278999, 264259, 52645080, 29331822, 29331824, 29331825, 29331826, 29331822, 29331824, 29331825, 29331826, 29331826, 29331827, 2656970, 264508, 264509, 264906, 265007, 265017, 265018, 265006, 265007, 264682, 264682, 264683, 264684, 264288, 264686, 21906769, 265020, 265021, 265022, 52644150, 33657023, 23657109, 27466265, 33657349, 18108374, 35696423, 356526486, 50432113, 2279000, 264487, 2648518, 60432113, 2279000, 264487, 2648518, 60432113, 2279000, 264487, 2648518, 60432113, 2279000, 264487, 2648518, 60432113, 2279000, 264487, 264567, 264486
2181	85764930 (4361, 4362)	85764930 (4361, 4362) Novel Protein sim. GBank gij3024689jspjQ15542jT2D4_HUMAN - TRANSCRIPTION iniTiaTion FACTOR TFiID 100 KD SUBUNIT (TAFII-100) (TAFII100)		Kinase	29331827, 264369, 18108376, 264564
2182		87637731 (4363, 4364) Novel Protein sim. GBank gil5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan Lelshmania major]		UNCLASSIFIED	22276996, 22278997, 22278999, 264259, 29331822, 56182435, 264112, 264764, 264288, 21906767, 21906768, 21906769, 33657109, 18108376, 60170394, 22279000, 22279002
2183	85460649 (4365, 4366)	85460649 (4365, 4366) Novel Protein sim. GBank gij3973406jgbjAAC77482.1 - (U17129) unknown [Rhodococcus erythropolis]			264760
2184		87760690 (4367, 4368) Novel Protein sim. GBank gij3114713 (AF061346) - Edp1 protein [Mus musculus]		tní	29331822, 29331825, 29331826, 56182435, 265011, 264685, 264686, 21906768, 18108370, 264629, 264631, 264636, 264557
2185	87826463 (4369, 4370)	87826453 (4369, 4370) Novel Protein sim. GBank gi 5106956 gb AAD39906.1 AF11361 - (AF113615) FH1/FH2 domain-containing protein FHOS [Homo sapiens]			29331824, 264907, 66712502, 264757, 265019, 264288, 264692, 56526486
2186		87739227 (4371, 4372) Novel Protein sim. GBank gi 2864625 emb CAA16972 - (AL021811) putative protein [Arabidopsis thaliana]		ATPase_associated	ATPase_associated 264259, 29331822, 29331826, 29331826, 56182435, 264592, 55812038, 264760, 264766, 55811957, 33657023, 55811576, 56182323, 264563
2187	87388173 (4373, 4374)			UNCLASSIFIED	35696052, 264905, 264906, 264907, 264908, 264510, 264511, 265008, 264910, 264768, 265019, 264762, 264681, 264766, 264691, 264692, 2696917, 264692, 35696423, 264681, 264635, 264637, 18108388, 284568, 264486
2188	87771708 (4375, 4376)	87771708 (4375, 4376) Novel Protein sim. GBank gil5107816[gblAAD40129.1]AF14941 - (AF149413) contains similarity to histone deacetylases; Pfam PF00850, Score=13.3, E=5e-10, N=1 [Arabidopsis thaliana]		histone	18108398, 56994075, 264259, 29331824, 29331825, 66714117, 29331827, 264906, 29331830, 265018, 265020, 26279002 56182323, 264559, 22279000, 22279002
2189		85693573 (4377, 4378) Novel Protein sim. GBank gi]3452357 (AF075724) - unknown [Legionella pneumophila]	Contains protein domain (PF01596) - O-methyltransferase		22278996, 264259, 29331826, 21906754, 264369, 264288, 263967

2190	87639197 (4379, 4380)	Novel Protein sim. GBank gij132575 sp P29315 RINI RAT -		didococologic	222220000000000000000000000000000000000
		RIBONUCLEASE INHIBITOR			29331826, 265008, 264910, 60170831,
					55812038, 52644296, 265010, 265018,
1					264685, 264688, 56181562, 21906769,
2191		95198928 (4381, 4382) Novel Protein sim. GBank gil5327002 emblCAB46272.11 -			33093917, 265022, 60170394, 22279000
		(Y18503) XAP-5-like protein [Homo sapiens]			29331625, 29331626, 29331830, 264510, 264511, 264910, 264593, 264594, 264556
2102	_				264559
7		1 (1203 10 (4303, 4304) Novel Protein Sim. GBank	Contains protein domain (PF00169) -		264558
2103		94140073 (4286 4286) Wilder Barandini CAEEL - MIG-10 PROTEIN	PH domain		
3		Novel Protein Sim. GBank gi 5420389 emb CAB46680.1		UNCLASSIFIED	56181686, 29331825, 29331827, 264508
		(AJ243460) proteophosphoglycan [Leishmania major]			264909, 265008, 264592, 60432229, 264288
					264684, 264766, 35695917, 33657023.
	· · · · · · · · · · · · · · · · · · ·				60431602, 60431528, 55810764, 55811576,
					65274791, 35695855, 60431850, 56182323,
2194		21418714 (4387, 4388) Novel Protein sim. GBank gil2773341 (AF040954) - putative			264502
		protein phosphatase 1 nuclear targeting subunit [Rattus		•	7000
1	_	norvegicus]			
2185	_	88083023 (4389, 4390) Novel Protein sim. GBank gij2832763jembjCAA15685.11 -		UNCI ASSIFIED	22278006 22278000 25606052 266006
		(AL009191) /prediction=(method:; /prediction=(method::			144,050, 444,06039, 33090034, 4630006,
		/match≂(desc;; /match=(desc;; /motif=(desc; IDrosophila		<u> </u>	21900/34, 203011/, 33093917, 263021,
		melanogaster]			(050/22, 35095855
2198		95091631 (4391, 4392) Novel Protein sim. GBank gil5262487lemblCAB45699 11.			COLUMN OF CONTRACT STATES
		(AL080076) hypothetical protein [Homo sapiens]			301023/3, 33096286, 222/8997, 22278999, 364350 30331833 66314447 6043330
					29331827 35696052 29331828 264508
				45	52644045, 56182435, 264510, 265007.
				2	265008, 265009, 60433438, 55812038,
	_			2	265010, 265011, 264448, 264288, 264686
				2	264687, 52644229, 21906765, 21906766,
				2	21906767, 35695917, 265022, 264691,
				<u> </u>	33657023, 264693, 18108370, 18108376,
				r (35696423, 55811576, 65274791, 35695855,
2197	95073813 (4393, 4394)	95073813 (4393, 4394) Novel Protein sim. GBank		7 2	264368 264769 21908385 264768 264769 21906765 21906766
		gil4929567[gb AAD34044.1 AF15180 - (AF151807) CGI-49		2	21906767, 29148627, 55811957, 35696286
		protein [Homo sapiens]			265020, 22278998, 265021, 264259.
				<u>e</u>	33657023, 264693, 29331824, 35696052,
				2	29331828, 18108370, 35695855, 264113,
				2	265008, 264910, 60432229, 56182323,
				<u></u>	33657402, 264758, 83373044, 21906754,
		•		2 0	265018, 265019, 22279002, 264482, 264448,
2198	88060914 (4395, 4396)	88060914 (4395, 4396) Novel Protein sim. GBank gij3548787 (AC005622) -		UNCLASSIFIED	264565, 264288, 264369
7		R30953_1 [Homo sapiens]			
				-	

2199		88054355 (4397, 4398) Novel Protein sim. GBank gil2739372 (AC002505) - hvoothelical protein fArabidoosis thallana			264105, 264110, 264112, 264688, 55811957, 33657023, 264692, 263967, 20281071,
					56526486
2200		87405385 (4399, 4400) Novel Protein sim. GBank gij3043634[dbj BAA25481] - [AB011127] KIAA0555 protein [Homo sapiens]		struct	29331824, 264763, 264768
	94316872 (4401, 4402)	94316872 (4401, 4402) Novel Protein sim. GBank gij3913470jspjO57314lDHBX_ANAPL - PUTATIVE STEROID DEHYDROGENASE SPM2	Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase	dehydrogenase	29331824, 35696052, 264905, 264907, 33657402, 55811386, 265017, 265019, 264288, 21906788, 35695917, 265020, 285022, 33657023, 33657109, 27486261, 18108370, 35696423, 35695855, 264556, 83373044, 87168518, 60432113
		91672385 (4403, 4404) Novel Protein sim. GBank gil5262665 emb CAB45767.1 - (AL080186) hypothetical protein [Homo sapiens]		UNCLASSIFIED	264489, 264259, 29331824, 60432289, 35696052, 264905, 264909, 264592, 265017, 265018, 265019, 18108351, 264762, 264448, 264369, 264288, 264766, 21906765, 21906766, 264691, 264692, 33657109, 264634, 264636, 264555, 264639, 264558, 264559, 83373044, 18108385, 264404, 22279002, 264482
2203		87761832 (4405, 4406) Novel Protein sim. GBank gij1172845 sp P46629 RB25_RABIT - RAS-RELATED PROTEIN RAB-25	Contains protein domain (PF00071) - glycoprotein Ras family	glycoprotein	52646365, 56994075, 264259, 29331822, 29331826, 29331827, 29331828, 264910, 265010, 265011, 87168559, 265018, 265019, 264605, 264288, 21906769, 35695917, 33657023, 264692, 33657109, 35695763, 18108376, 264638, 22279000, 264566, 264567
		88088671 (4407, 4408) Novel Protein sim. GBank gij121036 sp P29348 GBT3_RAT Contains protein domair GUANINE NUCLEOTIDE-BINDING PROTEIN G(T), ALPHA-G-protein alpha subunit 3 SUBUNIT (GUSTDUCIN ALPHA-3 CHAIN)	121036 sp P29348 GBT3_RAT Contains protein domain (PF00503) - UNCLASSIFIED NDING PROTEIN G(T), ALPHA G-protein alpha subunit LPHA-3 CHAIN)	UNCLASSIFIED	
2205		Novel Protein sim. GBank gil4589480 dbj BAA76768.1 - (AB023141) KIAA0924 protein [Homo sapiens]	Contains protein domain (PF00096) - dna_rna_bind Zinc finger, C2H2 type	dna_rna_bind	18108394, 18108397, 56182575, 60432049, 264259, 29331822, 29331824, 29331825, 265007, 265008, 265009, 265010, 265010, 265010, 265011, 265011, 265011, 265011, 265011, 264680, 21906766, 21906769, 264680, 264681, 264691, 264634, 18108368, 55811576, 65274791, 264634, 18108381, 18108384, 60432113, 22279002, 264563, 264566
2206	20620008 (4411, 4412)	20620008 (4411, 4412)		UNCLASSIFIED	264591
		Novel Protein sim. GBank gi 4557753 ref NP_000372.1 pMID1 - midline 1 protein	Contains protein domain (PF00622) - SPRY domain		29331822, 56182181, 29331827, 35696052, 52644045, 265006, 265019, 56181562, 55811957, 265021, 33657023, 35695763, 35695855, 60170394, 60432113, 264566
2208	Ī				264906, 265019, 18108351, 21906769
2209	_	87800420 (4417, 4418) Novel Protein sim. GBank gij3986746 (AF105228) - tuftelin [Bos taurus]		struct	264112, 265009, 264691, 18108365, 18108374, 264634, 20281166

2210	57152407 (4419, 4420)	2210 [57152407 (4419, 4420) Novel Protein sim. GBank		kinase	264603
		gij728837 sp P39194 ALU7_HUMAN - !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!			
2211	87341720 (4421, 4422)	87341720 (4421, 4422) Novel Protein sim. GBank gif728837[sp[P39194]ALU7_HUMAN - IIII ALU SUBFAMILY SO WARNING FNTRY IIII		oucogene	264685, 264686, 18108365, 22279002, 264482
2212	91223924 (4423, 4424)	gij3776027[emb[CAA09214] - [Arabidopsis thaliana]	Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase	helicase	22278995, 22278997, 22278999, 264092, 264094, 29331822, 66714117, 29331826,
					29331828, 264907, 52644045, 265009,
				_	60170831, 21906754, 87168559, 265017,
					265019, 18108351, 264683, 18108354,
					264369, 264766, 264687, 52644229,
					21906765, 21906766, 21906767, 21906768,
					265021, 33657109, 18108370, 18108374,
					264638, 56182323, 18108384, 18108387,
					87168518, 264565
2213	91219309 (4425, 4426)	91219309 (4425, 4426) Novel Protein sim. GBank gil5420387 emb CAB46679.11 -			56182575, 22278996, 22278997, 35696052,
		(AJ243459) proteophosphoglycan [Leishmania major]			264905, 66712502, 264908, 264828,
					56182435, 264112, 265008, 60431735,
					60433438, 21906754, 265010, 265011,
					265017, 265018, 265019, 18108351, 264765.
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					265021, 264693, 264629, 263974, 263976,
					18108379, 55811576, 264556, 264637.
					264558, 83373044, 22279002, 264482,
					264483

2214	95361453 (4427, 4428)	2214 95361453 (4427, 4428) Novel Protein sim. GBank	Contains protein domain (PF00725) - dehydrogenase	dehydrogenase	264488, 52644507, 18108394, 56182575,	
					22778997, 22278998, 22278999, 264490, 60432049, 264259, 26245080, 29331822, 29147620, 29331824, 66714117, 29331825, 29147620, 29331824, 66714117, 29331825, 29331827, 35696052, 29331828, 20281100, 264509, 264907, 265126, 264908, 29331830, 52644045, 26512502, 264908, 29331830, 52644045, 265124298, 265009, 60170813, 264512, 264691, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 264687, 5264692, 21906765, 219067	
2215	95419206 (4429, 4430) ,	95419206 (4429, 4430) Novel Protein sim. GBank gil1947160 (AF000298) - weak similarity to collagens; glycine- and proline-rich [Caenorhabditis elegans]			264909, 265006, 264555, 264558, 87168518	
2216	87614046 (4431, 4432)	87614046 (4431, 4432) Novel Protein sim. GBank gi]1572802 (U70854) - similar to Enterococcus faecalis TRAB (GI:388268) [Caenorhabditis elegans]		UNCLASSIFIED	264693	
2217	80589404 (4433, 4434)	80589404 (4433, 4434) Novel Protein sim. GBank gi 5031707 ref NP_005503.1 pGARP - glycoprotein A repetitions predominant	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	glycoprotein	264288, 33657109, 264556	r
2218	85518254 (4435, 4436)	85518254 (4435, 4436) Novel Protein sim. GBank gij3878636jembjCAA88953j - (Z49128) similar to cAMP-dependant protein kinase; cDNA EST EMBL:100719 comes from this gene; cDNA EST yk46548.3 comes from this gene; cDNA EST yk46548.5 comes from this gene; cDNA EST yk492f4.3 comes from	Contains protein domain (PF00069) - Eukaryotic protein kinase domain		35696423, 264563	,
2219	87614048 (4437, 4438)	87614048 (4437, 4438) Novel Protein sim. GBank gi 1572802 (U70854) - similar to Enterococcus faecalis TRAB (GI:388268) [Caenorhabditis elegans]	Contains protein domain (PF01963) - TraB family		264682, 264683, 264688, 264689, 264693, 18108370, 18108376	

2220	95354165 (4439, 4440)	2220 95354165 (4439, 4440) Novel Protein sim. GBank gi 4507261 ref NP_003145.1 pSTAT - statherin			264488, 18108394, 18108395, 35696286, 264259, 264097, 60432289, 264509, 264905, 264906, 264906, 264906, 264907, 29331830, 264908, 264510, 264511, 265007, 264512, 264910, 265009, 264594, 60433356, 264591, 264001, 264003, 265019, 264605, 264760, 264762, 264764, 264764, 264769, 264769, 264769, 264769, 264769, 264769, 264769, 264769, 264687, 18108358, 264628, 264624, 18108362, 18108362, 264634, 264634, 264638, 18108358, 264638, 264634, 264586, 264486, 264567, 264638, 18108358, 264634, 264634, 264638, 264634, 264488, 264567
2221	88060927 (4441, 4442)	88060927 (4441, 4442) Novel Protein sim. GBank gi 3549154 (AC005625) - R27328_1 Homo sapiens			
2222	84425892 (4443, 4444)	_		UNCLASSIFIED	264908, 265020, 35695855
2223	95091649 (4445, 4446)			UNCLASSIFIED	265010, 264685, 264690, 264693, 264628, 263974, 263976, 55811576, 264555, 264638, 83373044, 264483
2224	87388515 (4447, 4448)	87388515 (4447, 4448) Novel Protein sim. GBank gi[3876005]emb[CAA84799] - (Z35719) cDNA EST EMBL:D67419 comes from this gene; cDNA EST EMBL:C13853 comes from this gene; cDNA EST EMBL:C11578 comes from this gene; cDNA EST yk234a7.3 comes from this gene; cDNA EST yk234a7.5 comes from this gene; cDNA EST	Contains protein domain (PF01958) - UNCLASSIFIED Domain of unknown function	UNCLASSIFIED	264259, 264509, 56182435, 265006, 265008, 265009, 265009, 264757, 21906754, 18108351, 264693, 18108374, 18108385
2225		85749484 (4449, 4450) Novel Protein slm. GBank gi 1255847 (U53338) - C05E11.1 gene product (Caenorhabditis elegans)		transport	22278994, 22278995, 22278999, 52644045, 264600, 265019, 21906765, 21906769
2226		86978953 (4451, 4452) Novel Protein sim. GBank gil4826524 emb CAB42852.1			264259, 29331822, 29331824, 29331825, 29331827, 264508, 264908, 265007, 264691, 264634, 264486
2227				UNCLASSIFIED	22278999, 265006, 265008, 18108354, 29148629, 29148784, 27486261, 18108374, 264637, 18108384
2228		91227337 (4455, 4456) Novel Protein sim. GBank gij606976 (U16800) - ribonucleoprotein (Xenopus laevis)	Contains protein domain (PF00076) - dna_ma_bind RNA recognilion molif. (a.k.a. RRM, RBD, or RNP domain)	dna_ma_bind	264091, 264092, 264094, 29331822, 29331825, 66714117, 264693, 263972, 264639, 83373044, 264563
2229	88060931 (4457, 4458)	88060931 (4457, 4458) Novel Protein sim. GBank gil3549155 (AC005625) - R27328_2 Homo sapiens]		UNCLASSIFIED	

	_	7		 -	_		
264488, 264768, 52644507, 264769, 21906765, 21906766, 21906767, 21906769, 22278996, 22278996, 22278996, 22278999, 22278999, 256921, 264259, 2564509, 264509, 264907, 18108370, 18108374, 35696423, 33569555, 26907, 264910, 26455, 33657402, 21906754, 18108387, 265010, 265018, 265019, 26501	264563	18108394, 56182575, 22278995, 35696286, 22278997, 22278997, 22278997, 22278995, 254259, 29331827, 35696052, 264907, 56182435, 265006, 265007, 265008, 264910, 264758, 55812038, 264603, 265018, 264369, 18108351, 264686, 264764, 264689, 21906765, 21906766, 21906767, 21906765, 21906767, 21906769, 29148829, 35695917, 264690, 52644150, 264691, 33657023, 264693, 18108370, 18108374, 55811576, 35696855, 2646897, 2646897, 2646897, 264691, 356916876, 264690, 264691, 3659234, 264691, 3659234, 264691, 3659234, 264691, 3659234, 264691, 3659234, 264691, 36469	264906, 33657402, 265018, 264288, 264686, 265020, 264635, 18108385	56182435, 264369, 264688, 21906765, 265020, 264693, 264556, 56526486	22278997, 264563	264569, 264687, 264769, 265022, 264259, 60432049, 264691, 29331826, 60432289, 20281149, 264906, 264907, 264511, 265008, 265009, 264634, 264635, 264636, 264556, 264557, 264558, 60433136, 264595, 264559, 264559, 2643231	264488, 264769, 21906765, 21906766, 21906767, 21906768, 21906765, 21906769, 22278995, 22278996, 22278996, 22278998, 265020, 265021, 22278999, 264259, 29331824, 29331827, 35695763, 18108376, 35695855, 265007, 60432229, 33657402, 60433356, 60433438, 83373044, 18108385, 21906754, 18108387, 60432113, 22279000, 265019, 22279002, 264482, 18108351,
UNCLASSIFIED	UNCLASSIFIED	transcriptlactor		kinase	CLASSIFIED		
					Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat	Contains protein domain (PF00011) - eph Hsp20/alpha crystallin family	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat
	88060937 (4461, 4462) Novel Protein sim. GBank gi]3549154 (AC005625) - R27328_1 [Homo sapiens]	87762581 (4463, 4464) Novel Protein sim. GBank gij5281316 gb AAD41476.1 AF13312 - (AF133124) transcription factor IIIC63 [Homo sapiens]	87755292 (4465, 4466) Novel Protein sim. GBank gil4249733 gb AAD13780 - (AF109377) IdIBp Mus musculus]	87771817 (4467, 4468) Novel Protein sim. GBank gi 1706559 sp P54352 EAS_DROME - ETHANOLAMINE KINASE (EASILY SHOCKED PROTEIN)	91012316 (4469, 4470) Novet Protein sim. GBank gil4972734[gb]AAD34762.11 - (AF132174) unknown [Drosophila melanogaster]	ooous ist (4471, 4472) Nover Protein sim. GBank gij1082675 pir] B53814 - p20 protein - human	91012318 (4473, 4474) Novel Protein sim. GBank gil4972734 gb AAD34762.1 - (AF132174) unknown [Drosophila melanogaster]
95342915 (4459, 4460)	88060937 (4461, 4462)		87755292 (4465, 4466)				
2230	2231	2232	2233	2234	2235	0000	2237

2238	2238 Q4QQBB57 (447E 447E)			
	(Oct in the company)	Contains protein domain (PF00286) -		264509, 264907, 264629, 264634, 264564
2239	B7798688 (4477, 4478)	viiai coat protein		
				29331825, 265009, 264369, 33657109,
2240	94121471 (4479, 4480) Novel Protein sim. GBank gil2982311 (AED51240) -	Contains protoin domain (DE00470)		181083/0, 18108374, 264557, 264559
	Orobable ublauting-right and property of the p	Unicallist protein domain (PF00179) - upiquitin	uninbian	264488, 65274572, 56182575, 35696286,
	ליים ויים יים ליים אינים אינים אינים אינים אינים אינים ויים יים אינים אי	Outquian-conjugating enzyme		22278997, 22278999, 264259, 29331827,
				35696052, 264508, 52644045, 56182435,
				264511, 265007, 265008, 265009, 60433356,
				60433438, 55812038, 21906754, 33657084,
				55811386, 265018, 265019, 18108351,
				264683, 264288, 264768, 264687, 264688,
				264769, 21906765, 21906768, 21906769,
				35695917, 265021, 265022, 60170615,
_				52644150, 33657023, 33657182, 33657349.
				35695763, 18108370, 35696423, 35695855,
2241	80091951 (4481 4482)			87168518, 22279000
22.43	04000075 (4400 4404)		UNCLASSIFIED	264693, 264629
7 7 7	91220073 (4463, 4464) NOVEL Protein Sim. GBank		synthase	22278995, 22278996, 22278997, 22278998
	gij2494312/splP70541 E2BG_RAT - TRANSLATION		_	264259, 29331822, 29331824, 29331826
	INITIATION FACTOR EIF-28 GAMMA SUBUNIT (EIF-28			29331827, 29331828, 264509, 265007
	GDP-GIP EXCHANGE FACTOR)			265009, 264596, 21906754, 265010, 265011
				265017, 265018, 265019, 264448, 264369.
				264288, 52644229, 21906765, 21906766,
				21906767, 21906768, 21906769, 265020,
				265021, 33657109, 27486262, 27486264,
				18108374, 35695855, 264634, 264637,
				56182323, 83373044, 56526486, 87168518,
2243	78902026 (4485, 4486)		INCIASSIEIED	285008
2244	85723527 (4487, 4488)		Ţ	00000
	to BZIP transcription factor [Caenorhabditis elegans]		CACCASSIFIED	204004
2245	lar to	Contains protein domain (PF00534) - UNCLASSIFIED	UNCLASSIFIED	52645156, 22278995, 22278996, 22278997
		Glycosyl transferases group 1		22278999, 29331822, 29331824, 29331827
				264907, 264512, 60433438, 264758
				21906754, 265011, 264603, 264764, 264687
				21906767, 21906768, 21906769, 55811957,
				265022, 264691, 264629, 35696423, 264638,
				18108387, 60432113, 22279000, 22279002,
				264566

2246	94848710 (4491, 4492)	2246 94848710 (4491, 4492) Novel Protein sim. GBank gil4996096 dbj BAA78326.1 - (AB028069) activator of S phase Kinase [Homo sapiens]	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins		65274572, 22278995, 35696286, 22278996, 22278997, 22278999, 264269, 35696052, 264106, 264006, 264006, 265007, 265008, 60433438, 33109954, 87168559, 265018, 265018, 264288, 21906765, 21906767, 21906768, 21906769, 55811957, 35695917, 265020, 265022, 27486264, 18108374, 65274791, 35695855, 60432113
2247		87862542 (4493, 4494) Novel Protein sim. GBank gil854065 emb CAA58337 - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	52645156, 52646365, 52645080, 35696052, 33656970, 52646317, 33657084, 265017, 21906768, 21906769, 35695917, 33657109, 52645129, 33657182, 27486261, 27486262, 33657349, 27486265, 18108387
2248		.1pHABP - hyaluronan-binding	Contains protein domain (PF00089) - cathepsin Trypsin	cathepsin	264488, 264259, 264907, 29331830, 264909, 265007, 265009, 264595, 21906754, 65274444, 264603, 265019, 264762, 264448, 264288, 21906766, 55811957, 265021, 264691, 18108374, 264634, 264635, 264636, 264555, 264638, 264557, 264586, 264486
2249	94685662 (4497, 4498) 79827508 (4499, 4500)	2249 94685662 (4497, 4498) Novel Protein sim. GBank gil4038461 (AF107772) - TcST11	Contains protein domain (PF00515) - leph TPR Domain UNC	CASSIFIED	264766, 264628, 264636, 264637 264908, 18108374
2251		87385863 (4501, 4502) Novel Protein sim. GBank gij3218467 embjCAA07090.1 - (AJ006529) putative phosphatase [Gallus gallus]		UNCLASSIFIED	264259, 35696052, 264508, 56182435, 265009, 264592, 264593, 264760, 264448, 264684, 264288, 264690, 264628, 55811576, 264555, 264556, 264557, 264558, 264559, 264566
2252	87735867 (4503, 4504)	2252 87735867 (4503, 4504) Novel Protein sim. GBank GR735867 (AF145316) vacuolar ATP synthase subunit D proton pump delta polypeptide [Homo sapiens]	(PF01813) -		264092, 264094, 264259, 29331822, 66714117, 29331828, 264102, 264103, 264104, 264105, 264109, 264112, 264511, 265007, 60433356, 265010, 18108351, 21906767, 21906768, 264691, 263974, 263977, 264486, 264567
2253	91010703 (4505, 4506)			UNCLASSIFIED	65274572, 265019

52	95320031 (4507, 4508)	95320031 (4507, 4508) Novel Protein sim. GBank gil4502847 refiNP_001271.1 pCiRB - cold inducible RNA- binding protein	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	dna_ma_bind	264569, 18108394, 18108396, 56182575, 56994075, 35696266, 22278999, 284094, 60432049, 284094, 60432049, 284094, 29331822, 29331824, 29331825, 29331826, 29331826, 29331826, 29331826, 284509, 284509, 284509, 284510, 286509, 264511, 265007, 265009, 26409, 284510, 265006, 264511, 265007, 265010, 265011, 87168559, 265017, 265019, 264449, 264764, 264286, 284692, 264017, 265019, 264480, 264764, 264691, 33657023, 264629, 18108370, 60431528, 263967, 26818370, 60431528, 263967, 264629, 18108370, 60431528, 263967, 264629, 18108370, 60431528, 263967, 264629, 18108370, 60431538, 264638, 56182323, 264556, 263981, 264638, 56182323, 264556, 264586, 264586, 264565, 264586, 264565, 264586, 264565, 264565, 264566, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264565, 264566, 264565, 264566, 264565, 264566, 264565, 264565, 264565, 264566, 264565, 264486,
2255	91010546 (4509, 4510) ,	91010546 (4509, 4510) Novel Protein sim. GBank gij5541865 emb CAB51072.1 - (AL096858) hypothetical protein [Homo sapiens]	Contains protein domain (PF00076) - struct RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	struct	264567, 18108391 65274572, 56182575, 22278997, 22278999, 264259, 29331822, 29331825, 29331826, 29331827, 29331828, 284508, 264905, 264906, 264907, 66712502, 264908, 56182435, 264510, 264511, 265008, 264593, 264595, 21906754, 33109954, 87168474, 265011, 265017, 265019, 264682, 264764, 264369, 264268, 264766, 264685, 264686, 21906769, 256020, 60170615, 52644150, 264690, 264692, 246693, 33657109, 33657349, 264632, 264636, 52644332, 56182323, 22279000, 22279002
2256	87020531 (4511, 4512)	87020531 (4511, 4512) Novel Protein sim. GBank gij3327174[dbj BAA31655] - (AB014580) KIAA0680 protein [Homo sapiens]		UNCLASSIFIED	264768, 264689, 18108374
2257	80088235 (4513, 4514)			UNCLASSIFIED	22278996, 22278999, 264681, 21906765, 21906768, 264567
2258	88090516 (4515, 4516)	88090516 (4515, 4516) Novel Protein sim. GBank gi 3025446 (AC004528) - R32184 2 [Homo sapiens]	Contains protein domain (PF00060) - misc_channel Ligand-gated ion channel	misc_channel	264908, 264592, 264764

18108396, 65274572, 56182575, 22278997, 22278998, 264259, 29331822, 29331827, 264805, 66712502, 264908, 264909, 56182435, 265007, 265008, 60432229, 3657084, 87168559, 18108351, 26448, 264683, 264288, 264596, 56181562, 265021, 60170615, 264430, 3657109, 60431528, 18108374, 52644332, 264882	56182575, 265020, 264905, 264906, 264908, 35696423, 264511, 264635, 55812038, 264758, 265018, 265019, 264605, 264760, 264583		56182575, 22278999, 29331822, 29331825, 60432289, 29331827, 35696052, 264508, 66712502, 52644045, 56182435, 265006, 265008, 265009, 60433356, 55812038, 265010, 265017, 265019, 264288, 264369, 21906765, 21906767, 55811957, 35695817, 52644150, 33657023, 33657109, 55811576, 65274791, 56182323	22278994, 22278997, 264907, 264828, 52644150, 18108381, 264693, 18108374	264686, 264488, 264768, 264769, 264691, 264508, 264905, 264509, 264906, 264907, 264908, 264909, 35695855, 264510, 264511, 264512, 265007, 265009, 264638, 264760, 264767, 264766, 264766, 264766, 264766	264689, 264910, 264764	22278995, 22278997, 22278999, 264259, 265006, 265007, 265009, 60433438, 21906754, 265010, 265011, 265017, 26448, 264683, 264288, 264689, 21906765, 21906768, 35695917, 265021, 18108374, 264638, 22279000, 22279002, 264566,
UNCLASSIFIED	oxidase	UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	ribosomalprot
	Contains protein domain (PF00090) - oxidase Thrombospondin type 1 domain					Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF01305) - ribosomalprot Ribosomal protein L15 amino terminal region
2259 95364155 (4517, 4518) Novel Protein sim. GBank gil4884140 emb CAB43278.1 - (AL050110) hypothetical protein [Homo sapiens]	88084119 (4519, 4520) Novel Protein sim. GBank gij3080663 (AC004614) - similar to f-spondin proteins AB006086 (PID:g2529225) [Homo sapiens]	88074157 (4521, 4522) Novel Protein sim. GBank gi]3334526jemb CAA16138] - (AL021306) predicted using FGENEH [Homo sapiens]	91639292 (4523, 4524) Novel Protein sim. GBank gil4877759[gb]AAD31421.1]AF12444 - (AF124440) MAGE tumor antigen D1 [Homo sapiens]	87602495 (4525, 4526) Novel Protein sim. GBank gij3341697 (AC003672) - hypothetical protein [Arabidopsis thaliana]	87756525 (4527, 4528) Novel Protein sim. GBank gi 1657601 (U66220) - unknown [Nannocystis exedens] ,	86918663 (4529, 4530) Novel Protein sim. GBank gil477072 pir A48018 - mucin 7 precursor, salivary - human	87773456 (4531, 4532) Novel Protein sim. GBank gij3150479 (AF067212) - partial CDS [Caenorhabditis elegans]
59 95364155 (4517, 4							2266 87773458 (4531, 45

2267	87395838 (4533, 4534)	2267 87395838 (4533, 4534) Novel Protein sim. GBank gij3560229jembjCAA20697.1 - (AL031530) hypothetical protein [Schizosaccharomyces pombe]		UNCLASSIFIED	35696286, 264259, 29331824, 29331825, 35696052, 29331828, 264905, 284509, 264907, 264908, 264512, 265009, 264910, 264593, 33657402, 265010, 264768, 264762, 264448, 284288, 264369, 264768, 52644229, 35695917, 264691, 33657023,
9300		No. 1 P. 10. 20. 11. 11. 11. 11. 11. 11. 11. 11. 11. 1			18108382, 33657109, 35696423, 264634, 18108381, 87168518, 264566
9077		osossoor (4335, 4330) nover Protein sim. Gbank gij728832 sp P39189 ALU2_HUMAN - IIII ALU SUBFAMILY SB WARNING ENTRY IIII		cadherin	264488, 264259, 264509, 264595, 265010, 265017, 264766, 18108385, 264486
2269		88177977 (4537, 4538) Novel Protein sim. GBank gi[103418 pir S17885 - TcD37 protein - fruit fly (Drosophila melanogaster)		UNCLASSIFIED	56182575, 60432049, 265007, 265009, 264591, 87168559, 264605, 18108351,
					21906764, 265020, 264629, 60431528, 264638, 18108385, 18108387, 60432113
2270	80410327 (4539, 4540)				264763
	91010392 (4541, 4542)			cyto450	264909, 56182435, 265008, 55812038,
					55811957, 33657023, 264693, 33657109, 55810764, 55811576, 56182323
2272	84208220 (4543, 4544)			UNCLASSIFIED	264905, 264908
	95014271 (4545, 4546)	95014271 (4545, 4546) Novel Protein sim. GBank gil4176370 (AC005058) - similar	Contains protein domain (PF00462) -		52645156, 22278996, 22278999, 60432049,
		to calcium-independent phospholipase A2; similar to AC004392 (PID:03367519) (Homo sapiens)	Glutaredoxin		264259, 29331822, 29331824, 29331825,
					265006, 264593, 60433438, 21906754.
					265018, 264689, 21906765, 21906766,
					21906767, 21906769, 265021, 265022.
					60170615, 264691, 33657023, 264693,
					33657109, 27486264, 18108376, 35696423,
					35695855, 264630, 52644332, 264558. 56182323, 22279002
2274	91640217 (4547, 4548)	91640217 (4547, 4548) Novel Protein sim. GBank gil1480112 lemb CAA67961 -	Contains protein domain (PF00538) - histone	histone	52645156, 22278997, 22278999, 52645080,
		(X99642) HP1-BP74 protein [Mus musculus]	linker histone H1 and H5 family		29331824, 29331825, 29331826, 29331827,
					29331828, 264905, 264908, 52644045,
					264511, 265008, 265009, 60170831, 264591,
					21906754, 33109954, 265011, 265018,
					18108351, 264448, 264288, 264684, 264766,
					21906765, 21906766, 21906767, 21906768,
					52644150, 264693, 18108364, 35695763,
					18108374, 35696423, 264634, 264557,
					284638,52644332,83373044,18108385, EEEDS48E 87168648 DDD70000
2275	88082501 (4549, 4550)	88082501 (4549, 4550) Novel Protein sim. GBank gij3165406 (AC004755) -	Contains protein domain (PF00122) - transport	transport	100000000000000000000000000000000000000
_	-	fos37502_2 [Homo sapiens]	E1-E2 ATPase		
977	[1128/44/ (4551, 4552)]			UNCLASSIFIED	264555, 264556

tein 22278999, 35696052, 265008, 265019, 264369, 265020, 265022, 55810764, 264404, 22279002	SSIFIED 18108394, 22278997, 29331826, 60433356, 60433438, 21906754, 265018, 33657023, 264630, 264630, 264646		265008 12100054 265010 265010 265020		29331826, 35696052, 264107, 264906, 264909, 52644045, 265006, 33657402	60433356, 264758, 265011, 265019, 264681,	264683, 264684, 264686, 21906765, 21906767 21906768 21906769 60170616	264690, 52644150, 18108362, 264692.	18108368, 18108374, 263978, 264631,	18108381, 264559, 18108385, 56526486,	1		29331827, 29331828, 29331830, 33657402,	04433438, 87168474, 265019, 18108351, 24408767, 24408769, 88844087, 22887022	42 1 3 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	263972, 55811576, 87168518, 20281169		22278997, 22278999, 29331822, 29331824,	56182181, 29331825, 29331827, 35696052,	29146499, 264905, 66712502, 264908,	265007, 265009, 60432229, 264593,	60431735, 60433356, 33109954, 33657084,	55811386, 87168474, 265010, 265011, 1266018, 265011, 1266018, 266018,	ZOUTO, ZOUTO, 20011130, ZO4063, ZO4369,	21906768, 29148627, 21906769, 55811957	265020, 265022, 33657182, 27486261,	18108370, 264628, 18108374, 55810764,	18108379, 55811576, 35696423, 35695855,	264630, 60431850, 263981, 18108382,	
(0560) - glycopro	UNCLASSIFIED	UNCLASSIFIED		0400) - transpor				,				UNCLASSIFIED					UNCLASSIFIED		-		-					_				-
Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat				Contains protein domain (PF00400) - transpos	vvo domain, G-beta repeat																									
2277 88084123 (4553, 4554) Novel Protein sim. GBank gil2880079 (AC004142) - similar to murine leucine-rich repeat protein; possible role in neural development by protein-protein interactions; 93% similarity to D49802 (PID:o1369906) Homo sapiens!	16 8	80419375 (4557, 4558) Novel Protein sim. GBank gij119714 spjP13983 EXTN_TOBAC - EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN)		95293048 (4561, 4562) Novel Protein sim. GBank gil4240299 db BAA74928.1 -		-					87602829 (4563 4564) Novel Protein sim GRank ni1522020 (162840)	nucleopoda o54 (Rattus norvenicus)	enorge control and a control a				95362386 (4565, 4568) Novel Protein sim. GBank	gi 2495729 sp Q92556 Y281_HUMAN - HYPOTHETICAL	FRUI EIN KIAAU281 (HA6725)											
88084123 (4553, 4554)	941330 <i>7</i> 9 (4555, 4556)	80419375 (4557, 4558)	94239723 (4559, 4560)	95293048 (4561, 4562)					*		37602829 (4563 4564)	(, a.a. (a.a.) a.a		-) 5362386 (4565, 4566)			-										•
2277				2281							2282						2283		_			_	_							

ED 56182575, 35696286, 56994075, 29331824, 29331825, 35696052, 56182435, 60433438, 55812038, 33109954, 87168474, 87168559, 265018, 18108351, 264769, 27900765, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906769, 65274620, 33657109, 2748254, 24629, 55810764, 558111576, 35695855, 56182323, 56526486, 87168518, 22279000, 264567	264488 18108397, 35696286, 264092, 264259, 29331822, 29331826, 264906, 264908, 264511, 264512, 265009, 264910, 18108351, 264764, 264369, 264588, 264685, 264631, 264637, 18108381, 56182323, 264659, 18108385, 264464, 264563, 264565	22278997, 22278998, 22278999, 264259, 29331822, 60432289, 29331828, 35696052, 265018, 264684, 264288, 264686, 21906765, 21906767, 21906768, 21906769, 265020, 265021, 264692, 33657109, 18108376, 356923, 35695855, 264634, 22279000, 222790002, 264563, 264486	ED 22278996, 60432289, 264682, 264683. 264689, 18108374 263974, 263978	or 264488, 65274572, 56182575, 22278997, 22278999, 264259, 29331822, 29331824, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331828, 35696052, 264907, 264908, 52644045, 56182435, 264112, 265006, 265007, 264910, 265009, 60433356, 33657402, 264595, 55812038, 21906754, 264288, 264766, 21906768, 21906768, 21906769, 26491957, 265020, 265021, 264691, 23657109, 55811576, 56182323, 60170394, 83373044, 18108385, 56526486, 264654
UNCLASSIFIED	slruct	nuclease	UNCLASSIFIED	franscriptfactor
	Contains protein domain (PF00560) - struct Leucine Rich Repeat	Contains protein domain (PF00929) - nuclease Exonuclease		
2292 94328834 (4583, 4584) Novel Protein sim. GBank gil4803672 emb CAB42643.1 - (AJ133769) nuclear transport receptor [Homo sapiens]	87759213 (4585, 4586) Novel Protein sim. GBank gij3252981 (AF068921) - Ras- binding protein SUR-8 [Mus musculus]	86693580 (4587, 4588) Novel Protein sim. GBank gi 2062680 (U88964) - HEM45 [Homo sapiens]		94321251 (4593, 4594) Novel Protein sim. GBank gi 5689501 dbj BAA83034.1
94328834 (4583, 4584)	87759213 (4585, 4586)		95312200 (4589, 4590) 80030781 (4591, 4592)	94321251 (4593, 4594) ,
2292			2295 2296	

60424179, 56181686, 22278995, 35696286, 22278996, 22278998, 22278999, 264490, 264259, 28331822, 29331824, 66714117, 60424269, 35698052, 29331824, 66714117, 60424269, 35698052, 29331824, 66714117, 26182435, 264510, 265006, 60433438, 21906754, 33109954, 55811386, 264686, 264682, 264686, 264682, 264686, 264686, 264683, 21906767, 29148629, 55811957, 29148784, 35695917, 265020, 4108362, 33657023, 18108354, 33657000, 60431628, 18108370, 60431528, 18108374, 55810764, 35698423, 36595855, 264634, 60431850, 18108386, 5618223, 264636, 83373044, 18108385, 60432113, 22279000, 264482, 264567, 264486		264488, 22278996, 22278999, 264259, 29331824, 66714117, 35696052, 264509.	264905, 264906, 264907, 264908, 264909.	265008, 264910, 265009, 264758, 265010,	18108351, 264764, 264766, 264768, 264769,	21906766, 21906767, 35695917, 265021,	264691, 33657023, 35695763, 18108370.	18108374, 35696423, 35695855, 264631,	264553 264563	264908, 264758, 265017, 21906765, 83373044, 264563	T	18108370, 18108385	264239, 50432049, 264907, 264909, 264910, 60432229, 33657402, 265011, 265018.	264762, 264448, 264769, 264637, 264638,	ED 264259, 29331824, 21906767, 33657182,		65274572, 22278996, 264908, 265006, 21906769, 264691, 264486
collagen	UNCLASSIFIED									struct	UNCLASSIFIED		אניטכנ		UNCLASSIFIED		
													Contains protein domain (PF00047) - struct				
2298 95312207 (4595, 4599) Novel Protein sim. GBank gij3875051 jembjCAB02849j - (Z81050) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D65056 comes from this gene; cDNA EST yk366b12.3 comes from this gene; cDNA EST yk366b12.5 comes from this gene; cDNA EST yk366b12.5		94124346 (4599, 4600) Novel Protein sim. GBank gil2443886 (AC002294) - Unknown protein (Arabidoosis thaliana)								91235725 (4601, 4602) Novel Protein sim. GBank gil2143637 pir 84505 - calcium-	88084141 (4603, 4604) Novel Protein sim. GBank gi[287497 (AC004144) -	R34001_1 [Homo sapiens]	Novel Protein Sim. GBank gij4884194 emb CAB43Z20.1 /Al 040048\ hunothedical protein [Homo canions]	Is read to the country of the countr	94840434 (4507 4508) Novel Protein sim GBank	gi[2494162]sp[Q10005]YRY1_CAEEL - HYPOTHETICAL 39.9 KD PROTEIN T15H9.1 IN CHROMOSOME II PRECURSOR	90935911 (4609, 4610) Novel Protein sim. GBank gil4972666 gb AAD34738.1 - (AF132150) unknown [Drosophila melanogaster]
95312207 (4595, 4596)	80193720 (4597, 4598)	94124346 (4599, 4600)								91235725 (4601, 4602)					_		
2298	2299	2300								2301	2302		2303		25		2305

264488, 22278995, 22278996, 35696286. 22278997, 22278998, 22278999, 264259. 29331822, 29331824, 29331825, 29331826, 29331827, 29331827, 29331826, 264906, 264907, 29331830, 264908, 264909, 265008, 265008, 265009, 3365402, 21906754, 8565842, 265010, 265011, 264001, 265017, 265018, 264004, 265019, 18108351, 26448, 26428, 264604, 265019, 18108351, 26448, 265022, 33657023, 264692, 18108370, 18108374, 18108374, 18108387, 26438, 26438, 26438, 264639, 264639, 264488, 264630, 264634, 265021, 265021, 265021, 265021, 265021, 265021, 265031, 265031, 264639, 264639, 264639, 264639, 264639, 264638, 264488	264828	35696286, 264259, 29331822, 29331824, 264112, 264512, 264757, 21906754, 264288, 264690, 27486264, 264631, 264634, 264404	18108397, 22278996, 22278997, 22278998, 22278999, 60432049, 29331822, 29331826, 60432289, 60433229, 60433256, 60433356, 60433438, 65274444, 265010, 264690, 264681, 26448, 265031, 26488, 21906766, 21906768, 265020, 264691, 264693, 264693, 65274791	264508	18108397, 56182575, 22278996, 56994075, 264259, 29331824, 29331827, 264508, 264907, 56182435, 264510, 264511, 265006, 264512, 265007, 265008, 265009, 60433438, 33109954, 265010, 265011, 264603, 265017, 18108351, 264762, 264683, 264288, 264386, 33657023, 20281149, 20281069, 264628, 264372, 55811576, 35696423, 20281071, 264632, 26436, 18108387, 87168518, 22279000, 264563, 264486
kinasereceptor	UNCLASSIFIED	synthase	glycoprotein	UNCLASSIFIED	transcriptfactor
Contains protein domain (PF00400) - kinasereceptor					Contains protein domain (PF00013) - transcriptfactor KH domain
Novel Protein sim. GBank gil4929565[gbjAAD34043.1]AF15180 - (AF151806) CGI-48 protein [Homo sapiens]		Novel Protein sim. GBank giļ4758732[ref]NP_004522.1 pMOCS - motybdenum cofactor synthesis 2	95357218 (4617, 4618) Novel Protein sim. GBank gij3878059jembjCAB17070j - (Z99942) cDNA EST EMBL:D73444 comes from this gene; cDNA EST EMBL:D70905 comes from this gene; cDNA EST EMBL:D72208 comes from this gene; cDNA EST EMBL:D75030 comes from this gene; cDNA EST EMBL:D75944 comes from this gene; cDNA EST		87721189 (4621, 4622) Novel Protein sim. GBank gi[2137337]pir 48281 - gene mCBP protein - mouse
2306 95334940 (4611, 4612) Novel Protein sim. GBank gi 4929565 gb AAD34043. protein [Homo sapiens]	79415283 (4613, 4614)		95357218 (4617, 4618) N (2 cd E: E: E:	79601668 (4619, 4620)	87721189 (4621, 4622) N
2306	2307	2308	2309	2310	2311

56182575, 56994075, 35696286, 22278996, 22278997, 22278997, 22278997, 22278999, 264259, 29331822, 29331824, 29331825, 29331826, 29331827, 52644045, 60432229, 60433356, 55812038, 33109954, 21906754, 87168474, 265018, 18108351, 264288, 5264229, 21906765, 21906766, 21906767, 21906768, 21906769, 265022, 60170615, 33657023, 27486261, 27486264, 35696423, 35695855, 18108385, 22279000, 22279002	263981	52644507, 52646365, 52646842, 22278996, 22278997, 22278999, 264259, 52645080. 29331822, 29331824, 60432289, 25645080. 29331822, 29331824, 60432289, 29331826, 22331827, 29331828, 35696052, 33658970, 5264045, 265018, 265019, 265011, 265018, 265019, 265011, 265018, 265019, 265017, 265018, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265056, 21906768, 3365739, 35695763, 18108374, 35696423, 35695885, 52844332, 22279000, 22278002, 264563, 264567	264905, 264906, 264767, 264768, 264693, 55811576, 264635, 56182323, 18108385	52644507, 52645156, 52646365, 22278994, 22278995, 22278996, 56994075, 35696286, 22278997, 22278999, 22278999, 264259, 29331822, 52645080, 29331824, 29331825, 29331822, 26331827, 35696052, 29331828, 2644045, 264906, 264907, 29331830, 52644045, 264906, 264907, 26331830, 52644045, 265019, 264413, 265019, 26441296, 81168474, 265019, 2644229, 21906764, 265019, 2644229, 21906764, 265020, 2644150, 33657023, 18108362, 18108376, 35694332, 33595855, 264631, 264556, 52644332, 33593244, 18108385, 18108387, 87168518, 60432113, 22279000, 264566, 264567
		UNCLASSIFIED	UNCLASSIFIED	glycoprotein
·				Contains protein domain (PF00789) - glycoprotein UBX domain
2312 87549681 (4623, 4624) Novel Protein sim. GBank gi[2911264 (AC002550) - Unknown gene product [Homo sapiens]	80042533 (4625, 4626) Novel Protein sim. GBank gi 3043626 db BAA25477 - (AB011123) KIAA0551 protein [Homo sapiens]	Novel Protein sim. GBank gil5596714 emb CAB51401.1 - (AL035398) dJ796117.2 (CGI-51) [Homo sapiens]		94312191 (4631, 4632) Novel Protein sim. GBank gil5531827[gb AAD4488.1] - (AF078856) р47 [Homo sapiens]
87549681 (4623, 4624)			80430119 (4629, 4630)	
2317	2313	2314	2315	2316

2317	2317 87020571 (4633, 4634)			UNCLASSIFIED	22278998, 60432049, 264910, 60432229, 264686, 264687, 264688, 264689, 264558,
2					18108385
	2316 (79939878 (4635), 4635)			UNCLASSIFIED	265006, 264910
2318	95101781 (4637, 4638) N	Novel Protein sim. GBank gi 5262613 emb CAB45746.1 -			264488, 264569, 18108396, 52646365,
	<u> </u>	(AL080155) hypothetical protein (Homo sapiens)			22278994, 22278995, 22278996, 56994075,
					35696286, 22278997, 22278998, 264259,
					52645080, 29331825, 29331826, 29331827,
					29331828, 29331830, 56182435, 60170831,
					60432229, 60431735, 33657402, 21906754,
					52644296, 87168474, 265011, 87168559.
					265017, 265018, 265019, 18108351, 264448,
_				િ	18108354, 264288, 264369, 52644229,
			1	€***; \$**	21906764, 21906765, 21906766, 21906767.
				,	21906768, 21906769, 265021, 265022,
					52644150, 33657023, 52645129, 33657109.
_					27486264, 33657349, 35695763, 18108370.
					18108376, 18108379, 35696423, 264558.
					83373044, 18108385, 56526486, 87168518,
					254564, 264565, 264566
2320	2320 91622426 (4639, 4640) Novel Protein sim GBank	Novel Protein sim. GBank		kinase	22278994, 60432049, 60432289, 29331827,
	ō.	gi 728837 sp P39194JALU7_HUMAN - IIII ALU SUBFAMILY			264511, 265008, 52646317, 265017, 265019.
		SQ WARNING ENTRY !!!			21906765, 18108372, 18108387, 22279002
2321	94320377 (4641, 4642)	94320377 (4641, 4642) Novel Protein sim. GBank gi 3873837 emb CAB02700 -		UNCLASSIFIED	264488, 264687, 18108394, 264689,
		(281029) Similarity to S.pombe hypothetical protein			21906765, 18108397, 18108398, 21906767,
_		C1D4.09C (SW:Q10154); cDNA EST EMBL:T00543 comes			21906768, 65274791, 22278995, 35695855,
	#	from this gene; cDNA EST EMBL: T01062 comes from this			22278998, 265021, 265022, 264510, 265006,
	<u> </u>	gene; cDNA EST EMBL:T01321 comes from this gene;			264511, 264512, 265008, 60170615, 264555,
	<u> </u>	cDNA EST EMBL: T02288 com			264636, 264556, 18108361, 264259,
					60432229, 33657023, 264557, 264558.
					264693, 60433356, 264559, 60433438,
					29331824, 18108365, 18108348, 18108384.
					29331825, 18108385, 33109954, 29331827,
		-			56526486, 29146499, 265011, 60432113.
					265017, 265018, 264508, 264563, 264482,
					264509, 18108351, 264448, 264907, 264682,
					18108370, 264683, 264908, 264288, 264909,
2322	87803165 (4643, 4644)	85.11-	Contains protein domain (PF00106) - dehydrogenase	dehydrogenase	22278996, 264907, 264511, 264757,
		\Box	short chain dehydrogenase		18108351, 264768, 264638

5, 4646) 9	2323 94840445 (4645, 4646) Novel Protein sim. GBank gi 2494162 sp O10005 yRY1_CAEEL - HYPOTHETICAL 39.9 KD PROTEIN T15H9.1 IN CHROMOSOME II PRECURSOR	Contains protein domain (PF00226) - eph DnaJ domain		22278994, 22278995, 22278997, 60432049, 264259, 29331822, 33658970, 264509, 26182435, 264511, 265008, 60433358, 60433438, 55812039, 33109954, 21906754, 85658542, 87168474, 265011, 87168559, 265017, 265019, 264687, 264687, 18106351, 24906765, 21906765, 21906765, 21906765, 21906765, 21906767, 21906365, 265611, 87168367, 256501, 23655021, 33657023, 18108362, 265611957, 21906765, 21906767, 21906768, 55811957, 21906769, 265011957, 21906768, 265011957, 216011957, 2165011957,
				27486262, 55811576, 264631, 264555, 83373044, 87168518, 60432113, 22279002, 264592, 264593, 265020
N S	88165074 (4649, 4650) Novel Protein sim. GBank gi[5419865 emb CAB46377.1 - (4649, 4650) (4649		associated	265020
§ 5 5	84390962 (4651, 4652) Novel Protein sim. GBank gij231885jspjP29981jCP4C_BLADI - CYTOCHROME P450 Cytochrome P450 4C1 (CYPIVC1)	omain (PF00067) -	cyto450	265006, 264759, 35695855, 56182323
S &	88081648 (4653, 4654) Novel Protein sim. GBank gil4240227 dbj BAA74892.1 - (AB020676) KIAA0869 protein [Homo sapiens]		UNCLASSIFIED	
§ 6	83388428 (4655, 4656) Novel Protein sim. GBank gi]1245105 (U46463) - glutamine repeat protein-1 [Mus musculus]		UNCLASSIFIED	264259, 264508, 264905, 264905, 264907, 264907, 264908, 265007, 264512, 264910, 264758, 265010, 264766, 264768, 264769, 264639
12 E	87604478 (4657, 4658) Novel Protein sim. GBank gil1169343 sp P42209 DIF6_MOUSE - DIFF6 PROTEIN	Contains protein domain (PF00735) - UNCLASSIFIED Cell division protein	UNCLASSIFIED	60433438, 264995, 265011, 264765, 264694, 264629, 264635, 264636, 264638, 56182323, 60432113, 264566
			UNCLASSIFIED	265017, 264685, 60432113, 264088
N E E	86990463 (4661, 4662) Novel Protein sim. GBank gij5679136jgbjAAD46874.1JAF16093 - (AF160934) BcDNA,LD14189 IDrosophila melanogaster]		transport	265009 265009
ž V	87784182 (4663, 4664) Novel Protein sim. GBank gi 2104452 emb CAB08779 - (Z95397) unknown [Schizosaccharomyces pombe]		ATPase_associated	ATPase_associated 35696286, 22278999, 29331824, 00424209, 265008, 265008, 265018, 264448, 264764, 21906765, 35695917, 35695955, 264636, 22279000, 264566
ZUUUS	98206958 (4665, 4666) Novel Protein sim. GBank gij3879985 emb CAA92691.1 - (Z68318) cDNA EST CEMSD62F comes from this gene; cDNA EST EMBL:C07930 comes from this gene; cDNA EST EMBL:C09493 comes from this gene; cDNA EST yk415e8.3 comes from this gene; cDNA EST comes from this gene; cDNA EST		UNCLASSIFIED	56182575, 56994075, 29331826, 29331828, 264107, 33657402, 87168559, 264683, 35695917, 265021, 33657023, 263976
3 2 <u>22013</u>	94319788 (4667, 4668) Novel Protein and Surie, Calank gilde6270(gbl/AAB52261.21-(197002) similar to acyl-CoA dehydrogenases and epoxide hydrolases; Pfam domain PF00441 (Acyl-CoA_dh). Score=57.4, E-value=1.7e-16, N=2; contains similarity to Pfam domain PF00702 (Hydrolase), Score=57.4, E-value=1e-13, N=1 [C	Contains protein domain (PF00441) - dehydrogenase Acyl-CoA dehydrogenase	dehydrogenase	56182575, 29331825, 21906768, 264636, 83373044

2335	80046103 (4669, 4670)	2335 80046103 (4669, 4670) Novel Protein sim. GBank gij3283350 (AF062378) - calmodulin-binding protein SH4 1 Mus musculus	Contains protein domain (PF00612) - struct	struct	18108351, 21906769, 264555
2336		95196121 (4671, 4672) Novel Protein sim. GBank gil1929056 emb CAA72805 - (Y12090) putative 3,4-dihydroxy-2-butanone kinase [Lycopersicon esculentum]		kinase	264907, 35695917, 18108379
2337	95345610 (4673, 4674)	95345810 (4673, 4674) Novel Protein sim. GBank gil4495063 emb CAB39181.1 - (Z85986) dJ108K11.3 (similar to yeast suppressor protein SRP40) [Homo sapiens]		UNCLASSIFIED	35696286, 22278999, 56182181, 29331825, 60424269, 56182435, 33657402, 55812038, 55811386, 265017, 265018, 265019, 21906768, 35695917, 264691, 33657023, 33657109, 263972, 35696423, 35695855, 60432113
2338	87634045 (4675, 4676)	2338 87634045 (4675, 4676) Novel Protein sim. GBank gi[2224689 dbj BAA20829 - (AB002372) KIAA0374 [Homo sapiens]	Contains protein domain (PF00323) - UNCLASSIFIED Mammalian defensin	UNCLASSIFIED	18108394, 29331822, 66714117, 60432289, 264906, 56182435, 265009, 60433438, 264596, 265010, 265019, 18108354, 264288, 264369, 55811957, 265021, 33657023, 263976, 55811576, 264632, 56182323, 264639
2339		85663319 (4677, 4678) Novel Protein sim. GBank gil3873550 emb CAA22127 - (AL033534) serine-rich protein [Schizosaccharomyces pombe]		UNCLASSIFIED	35696286, 264592, 264369, 264691, 264558
2340	90937716 (4679, 4680)				65274572, 22278994, 35696286, 22278997, 22278999, 264259, 29331822, 60432289, 29331826, 29331826, 29331830, 265009, 33657402, 33109954, 265017, 265018, 264766, 264685, 21906769, 35695917, 264691, 264692, 35696423, 87168518, 22279000
2341	87775281 (4681, 4682)				264259, 264908, 264909, 264682, 22279000
2342		95334968 (4683, 4684) Novel Protein sim. GBank gil3874563jemblCAB027971- (Z81042) similar to Yeast hypothetical protein YEY6 like; cDNA EST yk206h5.3 comes from this gene; cDNA EST yk206h5.5 comes from this gene; cDNA EST yk303h1.3 comes from this gene; cDNA EST yk303h1.5 comes from this gene; cDNA	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat	kinase	264488, 65274572, 22278995, 22278996, 22278997, 22278998, 22278999, 60432049, 264259, 29331824, 29331824, 29331825, 66714117, 29331826, 60432289, 35696052, 264905, 264509, 29331830, 265006, 60170831, 60432229, 60433356, 87168474, 265017, 265018, 265019, 284448, 264369, 264288, 21906765, 21906766, 55811957, 35695917, 265020, 265022, 52644150, 33657023, 65274620, 33657109, 18108370, 18108378, 55810764, 35696423, 55811576, 264556, 264558, 18108385, 60432113, 264556, 264567, 26457, 26457, 264567, 26457, 26477,
2343		87775448 (4685, 4686) Novel Protein sim. GBank gi 4929741 gb AAD34131.1 AF15189 - (AF151894) CGI-136 protein [Homo sapiens]		UNCLASSIFIED	264901, 264512, 265011, 264663
2344				UNCLASSIFIED	264758
2345		94319799 (4689, 4690) Novel Protein sim. GBank gi[2506307]sp[P13944 CA1C_CHICK - COLLAGEN ALPHA [1(XII) CHAIN PRECURSOR (FIBROCHIMERIN)	Contains protein domain (PF00092) - collagen von Wiltebrand factor type A domain	collagen	264488, 264259, 66712502, 264759, 83373044, 264566

	22278997, 264511, 264683, 264684, 264768, 264688, 264691, 264691, 264692, 55811576	kinase 18108394, 35696286, 264259, 35696052, 264508, 264905, 264905, 264906, 264907, 264905, 264905, 264906, 264907, 264908, 264510, 264511, 265006, 265007, 264512, 265008, 265009, 264511, 265006, 264591, 265008, 265009, 264910, 264591, 264592, 264593, 264593, 264591, 264601, 264762, 18108351, 264764, 264288, 264766, 264768, 264689, 35695917, 264693, 264628, 18108370, 264629, 18108374, 35695423, 264631, 264631, 264507, 264638, 264631, 264486	35696052, 29146499, 264909, 264369			- UNCLASSIFIED) - UNCLASSIFIED 29331824, 264908, 265006, 265008
255411 (U53153) - one short Contains protein domain (PF00515) - proteaseinhib cerevisiae protease A inhibitor TPR Domain nort region of weak similarity ssion mediator protein elegans]						Contains protein domain (PF00076) - UNCLASSIFIED RNA recognition motif. (a.k.a. RRM, (RBD, or RNP domain)	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00076) - UNCLASSIFIED DNA recognition moif (a k a DBM
2346 94131820 (4691, 4692) Novel Protein sim. GBank gil1255411 (U53153) - one short Contains prot region of weak similarity to S. cerevisiae protease A inhibitor TPR Domain 3 (SP-P01099) and another short region of weak similarity to S. cerevisiae glucose repression mediator protein (SP:P14922) [Caenorhabditis elegans]		95196133 (4695, 4696) Novel Protein sim. GBank gi[1929056 emb CAA72805 - (Y12090) putative 3,4-dihydroxy-2-butanone kinase [Lycopersicon esculentum]	87776502 (4697, 4698) Novel Protein sim. GBank gil4884106jemb CAB43254.1 - (AL050062) hypothetical protein [Homo sapiens]		86968042 (4701, 4702) Novel Protein sim. GBank gil728832 sp P39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!	87337196 (4703, 4704) Novel Protein sim. GBank gil731637 sp P38760 YHH5_YEAST - HYPOTHETICAL RNA recognition motif 75.9 KD PROTEIN IN SPO13-ARG4 INTERGENIC REGION RBD, or RNP domain)	2353 91638784 (4705, 4706) Novel Protein sim. GBank gi[1346955]sp P48809 RB27_DROME - HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN 27C (HNRNP 48) (HRP48.1)	87337199 (4707, 4708) Novel Protein sim. GBank
94131820 (4691, 4692)	85330367 (4693, 4694)	95196133 (4695, 4696)	87776502 (4697, 4698)	88260594 (4699, 4700)	86968042 (4701, 4702)	87337196 (4703, 4704)	91638784 (4705, 4706)	87337199 (4707, 4708)
2346	2347	2348	2349	2350	2351	2352	2353	2354

7355 91538/85 (4709, 4710) Novel Protein sim. GBank gi (AL078455) hnRNP-like prot	Novel (AL078	Vovel Protein sim. GBank giļ4938503 emb CAB43861.1 - AL078465) hnRNP-like protein [Arabidopsis thaliana]	Contains protein domain (PF00076) - dna_rna_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	dna_rna_bind	56994075, 22278996, 35696286, 22278999, 264259, 29331826, 29331828, 29331828, 29331828, 29331828, 29348408, 264008, 26
					87168474, 265010, 265030, 203000, 201700, 87168474, 265010, 265030, 265080, 264691, 264692, 263967, 18108370, 87168518, 26279000
95327688 (4711, 4712) Novel Protein sim. GBank gil5138920 (AF092135) PTD014 [Homo sapiens]		5138920 gb AAD40377.1 - sapiens]			52644507, 22278995, 35696286, 22278996, 22278997, 22278998, 22278999, 264259, 29331824, 66714117, 29331825, 60432289, 35696052, 29331828, 264908, 66712502, 264512, 265007, 265008, 60170831, 60432229, 60433356, 60433438, 264758, 52646317, 33109954, 21906754, 55811386,
					87168474, 265017, 265018, 264605, 265019, 264681, 264682, 264448, 264369, 264288, 264886, 264788, 21906765, 21906766, 21906767, 21906762, 21906769, 265021, 60170615, 33657109, 27486264, 35695763, 55810764, 18108379, 35696423, 83373044, 18108385, 56526486, 264404, 60432113,
87775458 (4713, 4714) Novel Protein sim. GBank	Novel Protein sim. GBank			UNCLASSIFIED	22279000, 264482, 264563, 264566, 264486, 264567 264567 264488 264769 18108394 264259
		(AF151894) CGI-136			28331822, 18108370, 18108374, 264259, 29331822, 18108370, 18108374, 264510, 265017, 264462, 264563, 264762, 264565, 264566, 264389, 18108354
		4218005 (AC006135) - putative in-like) [Arabidopsis thaliana]		UNCLASSIFIED	22278997, 22278999, 264509, 264905, 264592, 18108351, 264681, 264682, 264769, 32833886, 18108374, 264556, 18108385, 264482
sin sim. GBank gil ins cDNA yk20f8.5 g1.5; coded for by y C. elegans cDN iNA yk20f8.5; cod		1086830 (U41264) - coded for S; coded for by C. elegans C. elegans C. elegans CA yk12b7.5; A yk36g6.5; coded for by C. elegans cDNA		UNCLASSIFIED	35696286, 22278938, 264905, 264511, 265007, 265008, 60433438, 264288, 264686, 21906769, 265020, 264692, 35695855, 264558, 56526486, 264563
2360 80046125 (4719, 4720) Novel Protein sim. GBank gij3881545 emb C (269804) cDNA EST yk428d5.3 comes from cDNA EST yk428d5.5 comes from [Caenorhabditis elegans]	yk log tz Novel Protein sim. GBank gij3B81545 (Z69904) cDNA EST yk42805.3 come cDNA EST yk42805.5 comes from thi [Caenorhabditis elegans]	3881545 emb CAA93779 - 5.3 comes from this gene; from this gene		UNCLASSIFIED	22278997, 29331826, 263981, 22279000
94232191 (4721, 4722) Novel Protein sim. GBank gi 746487 definition line found [Caenorhabditis e	Novel Protein sim. GBank gil746487 definition line found [Caenorhabditis e	746487 (U23514) - No nabditis elegans]			22278995, 22278999, 264512, 265009, 264757, 21906765, 65274620, 18108370, 60431528, 18108374, 264635, 60170394, 264482

2382	191721193 (4723 4724)	2382 01721193 (4723 4724) [Novel Protein eim GBank		UNCI ASSIFIED	22278999, 264259, 29331822, 29331824
		Š			60432289, 284509, 264512, 60432229,
		CHAIN IB (MYOSIN HEAVY CHAIN IL)			60433356, 264448, 264682, 264683, 264369.
					21906765, 21906768, 21906769, 60432113, 22279000, 22279002
2363	+	95006635 (4725, 4726) Novel Protein sim. GBank gil854065 emb CAA58337 -		UNCLASSIFIED	264907, 264629, 264635
2364		94827104 (4727 4728) Novel Protein sim GBank	Contains protein domain (PE00389) - reductase	reductase	264488, 18108394, 264887, 18108398,
		qij5639830 qb AAD45886.1 AF14601 - (AF146018)	D-isomer specific 2-hydroxyacid		22278996, 56994075, 35696286, 22278997,
	-	hydroxypyruvate reductase [Homo sapiens]	dehydrogenases		22278998, 264259, 66714117, 29331825,
					35696052, 264509, 264905, 264906, 264907.
					264908, 66712502, 264909, 264511, 265006.
					264512, 265007, 265008, 33657402, 264758,
					21906754, 87168474, 265010, 87168559,
					264603, 265017, 265018, 265019, 264760,
					264762, 18108351, 264448, 264764, 264683.
					264684, 264288, 18108355, 264766,
					18108358, 264689, 18108359, 21906765,
					21906766, 21906767, 35695917, 265020,
					265021, 265022, 60170615, 52644150,
					264691, 33657023, 264692, 18108364,
					33657109, 18108368, 18108370, 18108374,
					35696423, 35695855, 264635, 264556.
					264557, 264639, 60170394, 83373044,
					18108383, 18108384, 18108385, 18108388,
					56526486, 264482, 264564, 264486
2365		94140746 (4729, 4730) Novel Protein sim. GBank gil1840045 (U49082) -		transport	22278996, 22278998, 22278999, 264907.
		transporter protein [Homo sapiens]			264909, 264910, 33657402, 264758, 264600,
					264766, 264687, 264689, 21906765,
					21906767, 21906768, 21906769, 265021,
					33657023, 33657109, 83373044, 264566
2366	94312388 (4731, 4732)			UNCLASSIFIED	52644507, 264259, 29331824, 29331825,
					29331826, 29331827, 29331828, 264907,
					29331830, 264909, 264511, 265008,
					33657402, 264595, 52646317, 265017,
					265018, 265019, 264605, 264685, 264766,
					264689, 21906766, 21906769, 35695917,
					265020, 265021, 265022, 52644150,
					35695855, 52644332, 18108385, 18108387,
					264564, 264566
2367	94140910 (4733, 4734)	94140910 (4733, 4734) Novel Protein sim. GBank gij1065457 (U40410) - C54G7.4	Contains protein domain (PF00400) - UNCLASSIFIED	UNCLASSIFIED	35696286, 21906768, 55810764, 65274791, 264567
	_	gene product (Caenomandins eregans)	VVD COMBIN, G-Dela repeat		20,000
2368	94322190 (4735, 4736)				204020

2369	94314334 (4737, 4738)	2369 94314334 (4737, 4738) Novel Protein sim. GBank gil5360901 dbj BAA82158.1 -		struct	52644507, 52646842, 35696286, 264092.
	_	(ABUZ8343) a-nelix colled-coll rod nomologue (Homo sapiens			264094, 52645080, 35696052, 264107,
					28531930, 32044043, 283000, 283007, 285009, 52644298, 52644229, 264689
					21906765, 21906766, 35695917, 265020
					52644150 263967 33657109 27486265
					36606763 48408370 363074 48408334
					18108376 52644333 263084 18108374,
2370	(79804120 (4739, 4740)			INCIASSIFIED	264508 264000 264506
2371	г			T	264360
2372	Т-			Т	262067 262084
2373	Т	87418611 (4745, 4746) Novel Protein sim. GBank oil4589582ldbilBAA76813 11.		UNCLASSIFIED	2033167, 203301
		(AB023186) KIAA0969 protein [Homo sapiens]			29331829, 203010, 203019, 33093917, 264634, 60432113
2374		94123665 (4747, 4748) Novel Protein sim. GBank gil5105131 dbj BAA80445.1 -	Contains protein domain (PF01138) - UNCLASSIFIED		265006, 265007, 265008, 265009, 265011,
		(AP000061) 246aa long hypothetical ribonuclease PH	3' exoribonuclease family		264766, 35695917, 35695855, 263981,
2375		87731355 (4740 4750) Mayol Descrip cim Cont		T	264557, 264565
3		0	,	UNCLASSIFIED	60432049, 29331824, 264907, 52644045,
		9413311195pr477393KrB_MOUSE - SIGNAL RECOGNITION PARTICLE RECEPTOR BETA SLIBLINIT			264512, 60433356, 21906754, 52644296,
		ACD DETAIL			01 100333, 204440, 21900/03, 21900/00,
		(SN-BELX)	•		21906769, 33657023, 18108368, 55811576,
2376		87613744 (4761 4769) Novel Brotein sim CBank -19645435 (ACADETOR			52644332
2		Novel Frotein Sim. Grank gijz645435 (AFUU / /80) - CHD3	Contains protein domain (PF00628) - ATPase_associated	ATPase_associated	264259, 29331830, 264909, 264910, 265009, {
		Urosopniia meianogaster	PHD-finger		60433438, 21906754, 265017, 265018,
					265019, 264682, 264288, 264685, 21906767.
12.55	_			Ī	263972, 35695855, 87168518, 60432113
7757	_	93319669 (4/33, 4/34) Novel Protein sim. GBank gij5257005 gb AAD41239.1 -	Contains protein domain (PF01388) - UNCLASSIFIED		18108394, 65274572, 22278997, 22278999.
		(AF083249) Rb binding protein homolog [Homo sapiens]	ARID DNA binding domain		264095, 29331822, 29147620, 29331824,
					66714117, 29331825, 29331826, 29331828,
	_				33656970, 29146498, 29146499, 264509,
					265006, 265007, 265008, 265009, 60170831,
					265010, 265011, 265018, 55811150.
					18108351, 264764, 264288, 21906767,
					21906768, 29148627, 29148629, 265021,
					33657023, 33657109, 18108370, 18108374,
					18108379, 35696423, 264556, 83373044,
					18108385, 18108388, 56526486, 22279000.
22.20					22279002, 264563
62/0		9413/034 (4/35, 4/35) Novel Prolein Sim. GBank gi[1072198 (U40942) - No		UNCLASSIFIED	65274572, 56182575, 35696286, 264259,
		definition line found [Caenorhabditis elegans]			29331822, 29331824, 66714117, 29331825,
					29331826, 60432289, 29331827, 56182435,
					264510, 265009, 60433356, 87168474,
					265011, 265018, 264288, 21906765,
					33657023, 264557, 56182323, 83373044,
2379	65444324 (4757 475B)	65444324 (4757 4758) Novel Protein sim CBank 417227267 (AC004481)	(000 00 d) = 1 = 1 = 1 = 1 = 1 = 1 = 1 = 1		10100303, 22273002, 204402
3	(0011, 1011, 1001, 100)		Contains protein domain (Pr-00400) - kinase		265017, 264288, 21906768
			יים מיייניי, כ פטיני יבףכטי		

	86923062 (4759, 4760)		Contains protein domain (PF01410) - collagen Fibriliar collagen C-terminal domain	collagen	264908, 264910, 265011
		87608241 (4761, 4762) Novel Protein sim. GBank gil4455609 emb CAB36555 - (AL031846) dJ742C19.5 (novel Chromobox protein) [Homo sapiens]		helicase	56182575, 264091, 264093, 264259, 29331825, 264105, 264906, 60433356, 21906754, 265017, 265019, 264683, 264288, 264685, 264686, 264687, 264691, 264692, 264693, 55811576, 264636, 264567
2382		91225982 (4763, 4764) Novel Protein sim. GBank gil4325130 gb AAD17276 - (AF119716) dMi-2 protein [Drosophila melanogaster]	Contains protein domain (PF00628) - transport PHO-finger	transport	29331824, 60432289, 264905, 264596, 21906754, 264769, 265022, 264693, 263967, 33657109, 264629, 264631, 264558, 83373044, 60432113, 264482
2383		87442841 (4765, 4766) Novel Protein sim. GBank gi 1902982 db BAA19005 - (D89049) lectin-like oxidized LDL receptor [Bos taurus]	Contains protein domain (PF00059) - glycoprotein Lectin C-type domain	glycoprotein	265009, 21906765, 21906766
2384	95354766 (4767, 4768)	95354766 (4767, 4768) Novel Protein sim. GBank gl[2462851 (AF016252) - Spinophilin [Rattus norvegicus]	Contains protein domain (PF00595) - struct PDZ domain (Also known as DHR or GLGF).	struct	264488, 52644507, 52645156, 52646365, 35696286, 22278999, 52645080, 29331824, 29331826, 35696052, 29331828, 264906,
					26428, 5264449, 252000, 265009, 23109954, 33657084, 52644296, 265011, 265018, 2644296, 265011, 265018, 2644229, 21906765, 21906768, 265020, 22644150, 33657023, 264693, 65274620, 22645129, 33657182, 27486261, 27486262, 27486264, 33657349, 27486265, 35695763, 18108374, 35695855, 264634,
					264559, 18108385, 22279000, 22279002
2385	95419485 (4769, 4770) ,			UNCLASSIFIED	264488, 52644507, 52645156, 264887, 52646365, 22278995, 22278996, 22278997, 22278999, 264259, 52645080, 29331822, 29331826, 35696052, 52644045, 265006, 265007, 265008, 265009, 264910, 6043229, 60433356, 52646317, 21906754, 265019, 26448, 264683, 264686, 264687, 26906765, 21906766, 21906769, 265021, 265022, 264691, 264692, 65274620, 33657109, 18108370, 264631, 5264633, 22279000, 22279002, 264631, 264633, 264633, 26263, 264631, 264633, 26279002, 22279000, 22279002, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 26279002, 22279000, 22279002, 264633, 2646
2386		94742649 (4771, 4772) Novel Protein sim. GBank gil4929699[gb]AAD34110.1]AF15187 - (AF151873) CGI-115 protein [Homo sapiens]		glycoprotein	264488 22278995, 22278996, 22278997, 264289, 29146498, 264112, 264511, 60170831, 6043229, 264595, 60433438, 87168474, 87168559, 264682, 21906765, 21906766, 21906767, 21906769, 29148629, 35695917, 265021, 264690, 33657109, 264628, 18108376, 83373044, 60432113, 22279000, 264564, 264566, 264487

264634	264595	264488, 22278998, 22278999, 264509, 264905, 264906, 264906, 264907, 264908, 264909, 264909, 265006, 264511, 264512, 264910, 264591, 21906754, 264601, 264604, 264761, 18108351, 2644604, 264288, 264766, 264769, 264693, 3569423, 264635, 264636, 264555, 83373044, 22279000, 264488	\$2644507, 56182575, 22278995, 35696286, 22278996, 22278996, 22278997, 22278999, 29331822, 29331825, 29331826, 35696052, 264905, 55644045, 265009, 264758, 264759, 33109954, 52644296, 85658542, 265011, 265017, 265018, 264605, 52644229, 26905765, 21906765, 21906767, 21906768, 21906769, 35695917, 52644150, 33657023, 33657109, 33657349, 35695423, 18108376, 18108376, 186108376, 186108376, 56182323, 60170394, 83373044,	263976	35696286, 35696052, 264508, 264905, 264509, 264906, 264501, 264511, 264512, 264910, 265009, 264510, 264512, 264910, 265009, 264511, 264512, 264604, 264762, 264484, 264764, 264369, 264766, 264769, 264769, 264689, 35695917, 264629, 18108374, 263978, 35696423, 35695855, 264631, 264634, 264635, 264636, 264686, 264638, 60170394, 264639, 264565, 264486
UNCLASSIFIED	UNCLASSIFIED	dna_rna_bind	polymerase	UNCLASSIFIED	nuclease
		Contains protein domain (PF00170) - dna_rna_bind b21P transcription factor	Contains protein domain (PF00476) - polymerase DNA polymerase family A		Contains protein domain (PF00560) - nuclease Leucine Rich Repeat
1()		95310650 (4777, 4778) Novel Protein sim. GBank gil4758058 ref NP_004372.1 pCREB - cAMP responsive element binding protein-like 1	2390 94320912 (4779, 4780) Novel Protein sim. GBank gil1644239 dbj BAA12223 - (D84103) mitochondrial DNA polymerase gamma [Homo sapiens]		2392 94245016 (4783, 4784) Novel Protein sim. GBank gil4240169 dbj BAA74863.1 - (AB020647) KIAA0840 protein [Homo sapiens]
14997990 (4773; 4774)	11424604 (4775, 4776)	95310650 (4777, 4778)	94320912 (4779, 4780)	80036194 (4781, 4782)	94245016 (4783, 4784)
2387	2388	2389	2380	2391	2382

2202	05302533 /4785 4786)	2303 Toggozgas (4785 4786) Novel Protein sim GRank	Contains protein domain (PF00466) - Iribosomalprot	ibosomalprot	18108392, 60424179, 264489, 18108394,
2024	(2011, 2011)	gil4506667trefiNP 000993.1lpRPLP - ribosomal protein,	Ribosomal protein L10		18108397, 22278995, 56994075, 35696286,
					22278996, 22278997, 22278999, 264093.
					60432049, 264259, 29331822, 29147620.
					20281099, 29331824, 29331825, 66714117,
					60432289, 29331826, 29331827, 29331828,
					35696052, 29146499, 264508, 264509,
					264905, 264907, 264908, 66712502.
					52644045, 264828, 264909, 56182435.
					264112, 264113, 264510, 265006, 264511.
					265007, 265008, 265009, 264910, 264591,
					264593, 60433356, 264595, 60433438,
					52646317, 33109954, 21906754, 55811386,
					265010, 265011, 265017, 265018, 265019,
					264681, 264762, 18108351, 264763, 264682.
_					264764, 264683, 264369, 264288, 18108354.
					264766, 264686, 264687, 264688, 264689,
					18108359, 21906765, 21906766, 21906767.
					21906768, 21906769, 55811957, 29148629.
					29148784, 35695917, 265021, 265022.
					33657023, 264692, 264693, 18108364.
					33657109, 18108368, 27486261, 27486262,
		-			33657349, 35695763, 18108370, 263972,
					264629, 18108374, 263977, 18108376.
					263978, 55810764, 35696423, 35695855,
_					264634, 60431850, 264555, 264637, 264557.
					263981, 264558, 18108381, 60170394,
2204	_	04323256 (4787 4788) Novel Protein sim GBank gil4159888 (AC004908) - zinc	Contains protein domain (PF00096) - dna_ma_bind	dna_rna_bind	35696286, 22278997, 22278998, 56182181,
5	_	finger protein from gene of uncertain exon structure; similar		-	35696052, 265006, 264592, 55811386,
					265010, 265011, 265017, 265019, 264448,
					264683, 264288, 21906765, 21906768.
					21906769, 55811957, 35695917, 33657023,
_					65274620, 33657182, 33657349, 35695763.
					18108374, 18108376, 55810764, 55811576,
					35696423, 60170394, 18108385, 264564,
					264566, 264567
2395		95287212 (4789, 4790) Novel Protein sim. GBank		dna_ma_bind	264259, 29331824, 264910, 264288, 265021,
		gi[5712756]gb AAD47636.1 AF16079 - (AF160798) calcium			83373044, 18108387, 264563, 264568
		Iransporter CaT1 [Rattus norvegicus]			

2396	95096700 (4791, 4792)	95096700 (4791, 4792) Novel Protein sim. GBank gil106322[pirl B34087 - hypothetical protein (L1H 3' region) - human	Contains protein domain (PF00560) - nuclease Leucine Rich Repeat	nuclease	52646365, 18108397, 56182575, 35696286, 22278997, 22278999, 60432049, 264259, 29331822, 29331824, 29331825, 29331826, 29331827, 29331828, 264106, 264907, 29331830, 66712502, 264101, 60170831, 264591, 33657402, 60433438, 55812038, 3310954, 21906754, 33657084, 87168474, 265017, 265018, 265021, 60170615, 264692, 33657023, 65574620, 52645129, 23657182, 27486262, 27486265, 264631, 264556, 5264332, 264558, 83373044, 18108388, 87168518, 22279002, 264482
					52644507, 52645156, 56182575, 264259, 29147620, 264905, 264907, 264908, 264909, 264910, 264758, 52644296, 264603, 264604, 264762, 264681, 264764, 18108357, 264769, 21906768, 264693, 264628, 264635, 264638, 264639, 264564
2398	88047689 (4795, 4796)	88047689 (4795, 4796) Novel Protein sim. GBank gi 3258609 (AC005178) - H53_GS1 [Homo sapiens]		UNCLASSIFIED	
2399	87738965 (4797, 4798) ,	Novel Protein sim. GBank gil786117 (L41834) - nuclear protein [Ensis minor]		UNCLASSIFIED	56994075, 22276997, 22276998, 22276999, 264259, 29331822, 29331824, 66714117, 60432289, 264906, 29331824, 66714117, 60432289, 264906, 29331830, 56182435, 264112, 264910, 33109954, 21906754, 264764, 264765, 21906765, 21906766, 21906767, 265020, 255022, 60170615, 33657927, 26502, 60170615, 33657023, 18108374, 264556, 60170994, 264558, 87168518, 22279000, 22279002, 264564, 264566, 264487
2400	91214116 (4799, 4800)	2400 91214116 (4799, 4800) Novel Protein sim. GBank gi 2352822 gb AAB69285.1 - (AF008945) glucose-6-phosphatase [Haplochromis nubilus]		phosphatase	21906766, 52646842, 56994075, 33657182, 27486262, 52644296, 265017

52644507, 52645156, 52644229, 264688, 21906764, 21906765, 52646365, 52646842, 21906766, 21906767, 21906768, 22278995, 35695917, 56994075, 35696286, 22278996, 22278997, 265020, 22278998, 22278997, 265020, 22278998, 22278997, 265020, 22278998, 22278997, 264259, 33657109, 52645129, 29331826, 33657109, 52645129, 29331826, 33659162, 34696052, 27486261, 27486262, 33659406, 3264905, 35696423, 35695653, 264406, 264905, 35696423, 35695655, 264432, 55812038, 52646317, 18108385, 52644326, 87168474, 265010, 87168559, 80432113, 265017, 265018, 265019, 264563, 264288	264907, 264908, 264909, 264566		se 35696286, 29331826, 35696052, 265008, 265018, 21906769, 264564		52646842, 22278994, 22278996, 35696286, 22278997, 22278998, 22278999, 264092, 264093, 60432049, 264259, 29331822, 29331824, 29331825, 29331827, 35696052, 29331824, 285007, 264106, 264906, 52644045, 265007, 265008, 265009, 264910, 265491, 265011, 265018, 265009, 264369, 266485, 264686, 21906768, 21906769, 256485, 264632, 264633, 264632, 264633, 264632, 264633, 264632, 264633, 264632, 264633, 264632, 264633, 264632, 264639, 22279000, 22279002, 264653		ED 264910, 265010, 264448, 264557
phosphatase			dehydrogena	UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED
-		Contains protein domain (PF00062) - C-type lysozyme/alpha-lactalbumin family	Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase		CAP-Gly domain (PF01302) - struct		
2401 91214118 (4801, 4802) Novel Protein sim. GBank gil2352822 gblAAB69285.1 - (AF008945) glucose-6-phosphatase [Haplochromis nubilus]	91221408 (4803, 4804) Novel Protein sim. GBank gil4689258 gb AAD27832.1 AF12185 - (AF121859) sorting nexin 9 [Homo sapiens]	94135432 (4805, 4806) Novel Protein sim. GBank gil4929575[gb]AAD34048.1 AF15181 - (AF151811) CGI-53 protein [Homo sapiens]	95312605 (4807, 4808) Novel Protein sim. GBank gi[2315796 (AF016685) - similar to short chain-type dehydrogenases [Caenorhabditis elegans]	94311851 (4809, 4810) Novel Protein sim. GBank gil464178 db BAA03581 - (D14853) polyprotein [Hepatitis C virus]	98094501 (4811, 4812) Novel Protein sim. GBank gi[2773363 (AF041382) - microtubule binding protein D-CLIP-190 [Drosophila melanogaster]		87391503 (4815, 4816) Novel Protein sim. GBank giļ423442 pir S33513 - gene Fif protein - mouse
91214118 (4801, 4802)	91221408 (4803, 4804)	94135432 (4805, 4806)					
2401	2402	2403	2404	2405	2406	2407	2408

2409 94741770 (4817, 4818) Novel Potein sim. CBank gle96282 plyACA6022 21 Contains protein domain (PF00080) 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017664180 1017678333 14821, 4822) 1017664180 1017678333 14821, 4823 1017678333 14821, 4823 1017678333 14823, 4824 1017678333 101767833 1017678333	22278995, 22278996, 22278997, 264097, 29331822, 29331824, 29331827, 29146498, 52644045, 60433438, 33657084, 87168474, 264760, 21906767, 29148627, 29148629, 52644150, 33657023, 263967, 20281069, 18108374, 20281071, 56182323, 83373044, 18108385, 87168518	22278998, 264259	29331824, 29331827, 29331828, 264764, 264369, 33657109, 56182323	22278998, 29331827, 264907, 265011, 265017, 265018, 265019, 18108351, 21906766, 265020, 33657109, 264559, 18108385	264488, 264259, 29331826, 264508, 264905, 264509, 264906, 264907, 264908, 264908, 264910, 264511, 264612, 265008, 265009, 264910, 265011, 264682, 264766, 264686, 264768, 264686, 264768, 264689, 265021, 33657023, 18108370, 264628, 35695855, 264632, 264636, 264567, 264486	52645156, 52646842, 52646365, 18108398, 56182575, 22278994, 22278995, 56994075, 22278996, 35696286, 22278995, 56994075, 22278996, 35696286, 22278995, 56994075, 22278999, 264259, 5695407, 22278998, 22331824, 29331824, 29331824, 29331824, 29331824, 29331827, 29331824, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264630, 60433356, 33657402, 264594, 264687, 29146674, 21906766, 21906764, 264689, 21906766, 21906767, 29148627, 21906765, 21906766, 21906776, 29148627, 21906769, 265020, 265021, 60170615, 33657023, 264693, 264629, 18108370, 60431528, 264629, 18108376, 55811624, 264638, 264638, 264638, 264638, 264558, 56182323, 2843332, 264638, 282279902
26_CAEEL - HYPOTHETICAL 6 IN CHROMOSOME III 19149662621gbjAAC48052.2 - 11y to Pfam domain: PF00646 (F- 14.3e-05, N=1 [Caenorhabdiiis 1913114713 (AF061346) - Edp1 19154103361gbjAAD43038.1 - 1915262705[emb]CAB45778.1 - 10tein [Homo sapiens]	UNCLASSIFIED		Tu .	dna_rna_bind	struct	ubiquilin
4818) Novel Protein sim. GBank gil176601 sppP45966 yN2G_CAEEL - HYPOTHETICAL 20.8 KD PROTEIN T09A5.6 IN CHROMOSOME III (U64849) Contains similarity to Pfam domain. PF00646 (F-box), Score=28.7, E-value=4.3e-05, N=1 [Caenorhabdilis elegans] (Novel Protein sim. GBank gil3114713 (AF061346) - Edp1 protein [Mus musculus] (AF106685) myelin gene expression factor 2 [Homo sapiens] (AL080214) hypothetical protein [Homo sapiens] (AL080214) hypothetical protein [Homo sapiens] (AL080214) hypothetical by the protein [Homo sapiens]		Contains protein domain (PF00080) - Copper/zinc superoxide dismutase (SODC)		Contains protein domain (PF00076) - RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00038) - Intermediate filament proteins	
4820) 4822) 4828)	78. 11.9:	Novel Protein sim. GBank gil4966262[gb]AAC48052.2] - (U64849) Contains similarity to Pfam domain: PF00646 (F- box), Score=28.7, E-value=4.3e-05, N=1 [Caenorhabdilis elegans]	Novel Protein sim. GBank gi 3114713 (AF061346) - Edp1 protein [Mus musculus]	Novel Protein sim. GBank gi 5410336 gb AAD43038.1 - (AF106685) myelin gene expression factor 2 [Homo sapiens]	Novel Protein sim. GBank gi 5262705 emb CAB45778.1 - (AL080214) hypothetical protein [Homo sapiens]	Novei Protein R kappa B - human binding protein R kappa B - human
2410 87604860 (4819, 2411 87534633 (4821, 2412 87778332 (4825, 2414 94312590 (4827,	9 94741770 (4817, 4818) [87604860 (4819, 4820)				94312590 (4827, 4828)

415	88089002 (4829, 4830)	2415 88089002 (4829, 4830) Novel Protein slm. GBank gi 423915 pir A45439 - myosin I	Contains protein domain (PF00063) - struct	struct	264259, 264908, 60433356, 33657402,
		heavy chain - rat	Myosin head (motor domain)		21906754, 265018, 264687, 264689,
					21906769, 55811957, 265021, 264690.
					264691, 33657023, 264693, 35696423,
					56182323, 56526486
2416	94118356 (4831, 4832)	94118356 (4831, 4832) Novel Protein sim. GBank gil3025445 (AC004528) - R32184 1 [Homo sapiens]			264638
2417	87733334 (4833, 4834)	87733334 (4833, 4834) Novel Protein sim. GBank gi 1084944 pir S54495 - hypothetical protein YPR021c - yeast (Saccharomyces cerevisiae)	Contains protein domain (PF00153) - Mitochondrial carrier proteins		264094, 29331822, 29331824, 29331827, 264369
2418	94234349 (4835, 4836)	94234349 (4835, 4836) Novel Protein sim. GBank gij1176572 sp P45895 YNA4_CAEEL - HYPOTHETICAL 91.0 KD PROTEIN PAR2.4 IN CHROMOSOME III	Contains protein domain (PF00411) - UNCLASSIFIED Ribosomal protein S11	UNCLASSIFIED	56994075, 264091, 264259, 29331824, 29331825, 60432289, 29331828, 264905, 264907, 264511, 265009, 60432229, 21006754, 8216856, 255019, 264682
					21906768, 21906769, 265020, 265021, 23657023, 65274620, 18108370, 56811576, 264534, 60170394, 18108385, 22279000, 22279002, 264568
2419	82374249 (4837, 4838)	82374249 (4837, 4838) Novel Protein sim. GBank gi 284006 pir S18732 - autoantigen, 64K - human		struct	264569, 264762, 264448, 264691, 264631, 264634, 264555, 264556, 264638, 264558
2420	94844244 (4839, 4840)	9484244 (4839, 4840) Novel Protein sim. GBank gij1076211 pir S50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardtii		ÜNCLASSIFIED	29331824, 29331825, 29331828, 60432229, 33109954, 85658542, 87168474, 265018, 264288, 265020, 264564
421				UNCLASSIFIED	264909, 264768, 264638
2422	7 -	88084714 (4843, 4844) Novel Protein sim. GBank gil2224567 dbj BAA20772 - (AB002311) KIAA0313 [Homo sapiens]	Contains protein domain (PF00617) - transport RasGEF domain	transport	18108392, 18108394, 18108398, 264906, 265006, 265010, 18108351, 18108374, 18108385
2423		ARDSR300 (4845, 4845) Novel Protein sim GBank	Contains protein domain (PF00069) - kinase	kinase	264259, 60432049, 29331822, 29331826,
3		nover Tokell all. Obasin gi 4505153 ref NP_002392.1 pMEKK - MAP/ERK kinase kinase 3	Eukaryotic protein kinase domain		60432289, 29331828, 265008, 265009, 60433356, 21906754, 265017, 265018, 265019, 21906766, 21906768, 21906769, 265020, 265021, 20281149, 263971, 60432113
424	94854047 (4847, 4848)	2424 94854047 (4847, 4848) Novei Protein sim. GBank gil2988398 (AC004381) - Unknown gene product [Homo sapiens]	·	UNCLASSIFIED	56182575, 35696286, 22278997, 60432049, 264259, 29331826, 29331828, 264905, 66712502, 29331830, 60433356, 265011, 265019, 264766, 21906768, 55811957, 264692, 33657023, 33657109, 55811576, 56182323, 83373044, 18108385, 18108388, 60432113, 22279000
2425		87415981 (4849, 4850) Novel Protein sim. GBank gil2077932 dbi BAA19879 - (1086556) Protein Kinase (Rattus norvedicus)	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264634
426	87613945 (4851, 4852)	2426 87613945 (4851, 4852) Novel Protein sim. GBank gil2039368lgb AAB53003.11- (U94619) circulating cathodic antigen [Schistosoma mansoni]		UNCLASSIFIED	22278996, 22278998, 264259, 264102, 264512, 265008, 21906767, 18108370, 18108374, 263976

17622693 (485	4.	1854)		Contains protein domain (PF00573) - ribosomalprot	ribosomalprot	264259, 20281099, 35696052, 265008,
		٠,	37.1JAF13296 - (AF132962) CGI-28	Ribosomal protein L4/L1 family		264594, 265011, 264760, 18108351, 264682,
			protein [Homo sapiens]			264683, 264369, 264684, 264686, 264687, 264689, 21906766, 264691, 264692,
						18108374, 18108377, 264557, 264639.
						18108385
35732889 (4855, 48	æ	<u>2</u>	85732889 (4855, 4856) Novel Protein sim. GBank gij1537070 (U63840) -			22278996, 22278999, 35696052, 21906754,
			nucleoporin p54 [Rattus norvegicus]			264288, 21906765, 21906768, 21906769,
7760776 (4857 48	19	1681	87750775 (1857 1858) Novel Destein eim CBank mighton (Modats)	Contains protein domain (PE00711) - HNCLASSIEIED	INCI ASSIFIED	22278999 29331824 264906 264909.
) + (100+) 0 (360) 10	•	5		Beta defensins		264511, 265009, 21906754, 265017, 265018,
						265019, 264448, 264683, 264288, 21906765.
						21906768, 265021, 264693, 18108381
86948827 (4859, 4860)		100			UNCLASSIFIED	264112, 264691
37649884 (4861, 4		862)	87649884 (4861, 4862) Novel Protein sim. GBank gij3860729 emb CAA14630	Contains protein domain (PF01728) -		29331826, 29331827, 35696052, 29146499,
•			(AJ235270) CELL DIVISION PROTEIN FTSJ (ftsJ)	FisJ cell division protein		264905, 264906, 264681, 264288, 264689,
			[Rickettsia prowazekii]			21906765, 264692, 35696423
90083033 (4863, 4		864)	80083033 (4863, 4864) Novel Protein sim. GBank gi 3876367 emb CAA93287 -	_	protease	264634, 264558
			(Z69360) Weak similarity to Elmeria thrombospondin (PIR			
		_	Acc. No. A45517); cDNA EST EMBL:M89266 comes from			
			this gene; cDNA EST yk295b9.5 comes from this gene			
			[Caenorhabditis elegans]			
80055092 (4865, 4		(998	80055092 (4865, 4866) Novel Protein sim. GBank gij2224593 dbj BAA20784 · · · · · · · · · · · · · · · · · · ·	Contains protein domain (PF00096) - Itranscriptfactor Zinc finger: C2H2 type	transcriptfactor	264569, 264905, 265018, 264762, 264683, 264691, 264556, 264557, 264639, 264558
19520148 (4867, 4868)		868)				264563
20759044 (4869, 4870)		970			UNCLASSIFIED	264555
38044008 (4871, 4		1872)	88044008 (4871, 4872) Novel Protein sim. GBank gil 1263289 (U47856) - fibroin-4		UNCLASSIFIED	265008, 264758, 265010, 264689, 27486251,
•			(Araneus diadematus)			263972, 18108374, 18108381
83363424 (4873,		1874)	83363424 (4873, 4874) Novel Protein sim. GBank gij3641352 (AF091234) - putative		UNCLASSIFIED	29331828, 265007, 265009, 265017, 264760,
			franscription factor (Mus musculus)			264685, 264693, 264565
94143473 (4875, 4		(9/8)	94143473 (4875, 4876) Novel Protein sim. GBank gil3860014 (AF091088) -	Contains protein domain (PF01256) - UNCLASSIFIED	UNCLASSIFIED	29331826, 264508, 264905, 264509, 264906,
			unknown [Homo sapiens]	Uncharacterized protein family		264907, 264908, 264909, 264511, 265006.
				UPF0031		204012, 204910, 200009, 204031, 30607402,
		•				Z19Ub/54, Z05U11, Z04/6U, Z04/04, Z04005,
						204000, 204700, 33093917, 33037023, 364603 364631 364632 56182323 364558
						83373044 264563 264564 264565 264566
						264567

60424179, 18108397, 56182575, 56181686, 56994075, 22278996, 35696286, 22278997, 22278999, 264259, 52645080, 29331822, 56182181, 29331824, 60424269, 66714117, 29331825, 60432289, 29331826, 29331827, 29331825, 6043229, 29331826, 29331827, 29331828, 29331827, 26331828, 264509, 264905, 264906, 52644045, 60431735, 33109954, 21806754, 33857084, 55811386, 52644296, 81168571, 265019, 18108351, 264448, 264288, 264768, 5265019, 18108351, 264488, 264768, 21806765, 21806764, 21906765, 2186526, 21806764, 23857182, 27486262, 33657182, 263977, 55810764, 3569423, 65274791, 3569565, 60431850, 56182323, 60432113, 22279000, 22279902, 264567	29331822, 29331824, 29331825, 29331827, 35696052, 264508, 264907, 264510, 265018, 265019, 26448, 264369, 265020, 265021, 56182323, 264639, 22279002	264488, 264629, 18108374, 264564	22278996, 22278999, 29331822, 264768, 264693	264906	264259, 18108382, 18108383, 18108385, 22279000	264259, 35696052, 264369, 18108361	265011, 264689, 33657023, 263981, 18108385	56182575, 264259, 29331824, 29331825, 29331827, 60433356, 60433438, 264758, 265018, 264692, 65274620, 60431528, 65274791, 56182323
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transcriptfactor	transcriptfactor		proteaseinhib	cadherin
Contains protein			Contains protein domain (PF00560) - UNCLASSIFIED Leucine Rich Repeat	1170658 sp Q02975 KID1_RAT Contains protein domain (PF01352) - Iranscriptfactor FACTOR KID-1 3 17)	Contains protein domain (PF00170) - transcriptfactor bZIP transcription factor		Contains protein domain (PF00095) - proteaseinhib WAP-type (Whey Acidic Protein) Your-disulfide core*	
2439 94850850 (4877, 4878) Novel Protein sim. GBank gil4263519[gb]AAD15345] - (AC004044) small nuclear riboprotein Sm-D1 [Arabidopsis thaliana]		87623914 (4881, 4882) Novel Protein sim. GBank gi 3024889 sp P56524 Y288_HUMAN - HYPOTHETICAL PROTEIN KIAA0288 (HA6116)	87273590 (4883, 4884) Novel Protein sim. GBank gi 4506013 ref NP_002703.1 pPPP1 - protein phosphatase 1. regulatory subunit 7	A S C C	88086345 (4887, 4888) Novel Protein sim. GBank gi 4758824[ref]NP_004280.1[pNRF3 - nuclear factor (erythroid-derived 2)-like 3	97338636 (4889, 4890) Novel Protein sim. GBank gij2135950 pir S58222 - PQ-rich protein - human	88059293 (4891, 4892) Novel Protein sim. GBank gil4753887 emb CAA05409.2 - (AJ002424) p65 protein [Rattus norvegicus]	94845149 (4893, 4894) Novel Protein sim. GBank gil4885613 ref NP_005409.1 pST5 - suppression of tumorigenicity 5
94850850 (4877, 4878)	2440 87641733 (4879, 4880)	87623914 (4881, 4882)	87273590 (4883, 4884)	84305949 (4885, 4886)	86086345 (4887, 4888)	87338636 (4889, 4890)	88059293 (4891, 4892)	94845149 (4893, 4894)
2439		2441	2442	2443	2444	2445	2446	2447

2448				UNCLASSIFIED	22278996, 22278997, 22278999, 29331826, 35696052, 264107, 264110, 87168474, 87168559, 18108351, 21906767, 21906769, 27486262, 263976
2449		87869075 (4897, 4898) Novel Protein sim. GBank gij728837 sp P39194 ALU7_HUMAN - !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!		cadherin	264259, 264828, 265007, 264595, 265021, 56526486
2450	_			UNCLASSIFIED	264906
2451	91014563 (4901, 4902)		Contains protein domain (PF00071) - glycoprotein Ras family		264093, 29331822, 29331824, 29331825, 66714117, 29331826, 29331828, 35696052, 264907, 66712502, 29331830, 264910, 265009, 264758, 265017, 265018, 264762, 26448, 264288, 21906767, 265021, 33557023, 264693, 33657109, 263969, 83373044, 18108385
2452	91230509 (4903, 4904)	91230509 (4903, 4904) Novel Protein sim. GBank gil1504034 dbj BAA13216 - (D86980) KIAA0227 [Homo sapiens]		ізотегаѕе	264102, 264112, 26468B, 263972, 18108374, 83373044, 264563
2453		84201088 (4905, 4906) Novel Protein sim. GBank gil2880079 (AC004142) - similar to murine leucine-rich repeat protein; possible role in neural development by protein-protein interactions; 93% similarity to D49802 (PID:g1369906) [Homo sapiens]	Contains protein domain (PF00560) - ngfrecep Leucine Rich Repeat		264509, 264512, 18108385
2455		IIRE PIOIEIN - MAIZE 95288301 (4909, 4910) Novel Prolein sim. GBank gijs43817 sp p35585 AP47_MOUSE - CLATHRIN COAT ASSEMBLY PROTEIN AP47 (CLATHRIN COAT ASSOCIATED PROTEIN 47 (GOLGI ADAPTOR AP-1 47 KD PROTEIN) (HA1 47 KD SUBUNIT) (CLATHRIN ASSEMBLY PROTEIN ASSEMBLY PROTEIN COMPLEX 1 MEDIUM CHAIN)	DZIP transcription factor Contains protein domain (PF00928) - glycoprotein Adaptor complexes medium subunit family		264509, 264905, 264907, 264908, 264509, 264909, 264510, 264511, 264512, 265009, 264910, 264511, 264512, 265009, 264910, 264501, 264758, 264760, 264760, 264761, 264603, 264760, 264760, 264761, 264602, 3365703, 264692, 3365703, 264692, 3365703, 264692, 3365703, 264693, 33557109, 264637, 264638, 264639, 33373044, 1810835, 264563, 264696, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264591, 264691, 264591, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264631, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264631, 264633, 264623, 264623, 264623, 264623, 264623, 264633, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264633, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264623, 264633, 264623, 264623, 264633, 264623, 264623, 264633, 264623, 264623, 264623, 264633, 264633, 264623, 264623, 264633, 264623, 264633, 264623, 264633, 264633, 264623, 264633, 264633, 264633, 264623, 264633, 264633, 264633, 264623, 264633, 264633, 264633, 264623, 26463
2456	88166700 (4911, 4912)	2456 88166700 (4911, 4912) Novel Protein sim. GBank gil2588630 (AC003079) - Ankyrin-like; 54% similar to 2022340A (NID:g1092123) in exons	2588630 (AC003079) - Ankyrin-Contains protein domain (PF00023) - kinase A (NID:g1092123) in exons Ank repeat		60170394, 18108385, 264563, 264564, 264566, 264567 264693
		spanning 43974 to 11551 of clone. [Homo sapiens]			

	UNCLASSIFIED 264691, 264693, 264634, 264559	ort 27486265		UNCLASSIFIED 264259, 29331828, 264910, 18108351, 18108370, 18108374	264909, 264758, 264684, 18108374, 264637, 18108385	4SSIFIED 264681, 264566	264489, 52646842, 22278995, 35696286, 22278996, 22278999, 22278999, 22278999, 264259, 29331822, 29331824, 66714117, 29331825, 29331822, 29331827, 35696052, 29331828, 264509, 52644045, 264510, 264511, 264512, 265008, 60170831, 264593, 52646317, 33109954, 33657084, 265017, 265018, 265019, 264762, 264448, 264764, 264288, 264766, 21906765, 21906766, 21906767, 21906768, 21906769, 265021, 33657023, 33657109, 18108370, 18108381, 60170394,
חאכד	חאכר	Contains protein domain (PF00097) - transport Zinc finger, C3HC4 type (RING finger)	kinase	חאכדי		Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF01399) - protease PCI domain
	85675304 (4915, 4916) Novel Protein sim. GBank gil2384942 (AF022985) - Similar to collagen [Caenorhabditis elegans]	1 AC00499 - (AC004997) AA538043 (NID:g2284036)			86998002 (4923, 4924) Novel Protein sim. GBank gil5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]		nsi
94118375 (4913, 4914)	85675304 (4915, 4916)	2459 87551913 (4917, 4918) 	94315289 (4919, 4920)	87645147 (4921, 4922)	86998002 (4923, 4924)		2464 91219957 (4927, 4928) 6

2465	95357483 (4929, 4930)	2465 95357483 (4929, 4930) Novel Protein sim. GBank	Contains protein domain (PF00069) - oncogene	о исовеле	18108392, 52644507, 52645156, 52646365,
9976	1007, 1007, 007,033	leukemla viral oncogene homolog 1			22278998. 264259, 29331822, 29331824, 29331825, 60424269, 60432289, 29331827, 35696052, 29331828, 264907, 29331827, 255644045, 264909, 56182435, 264511, 265007, 265009, 265190, 26510, 26501, 87168559, 264910, 33657402, 265010, 265011, 87168559, 264600, 265017, 265010, 265011, 87168559, 264600, 265017, 265018, 265019, 18108351, 264369, 265020, 265021, 264686, 264631, 60170394, 26526486, 87168518, 60432113, 264563, 264566, 264487
i	85681386 (4931, 4932)	85681386 (4931, 4932) Novel Protein sim. GBank gil4321619 gb AAD15788.1 - (AF051098) seven transmembrane domain orphan receptor [Mus musculus]			264369
2467	88059465 (4933, 4934)	Novel Protein sim. GBank gij3513300 (AC005595) - F16601_1, partial CDS [Homo sapiens]		UNCLASSIFIED	56994075, 264908, 21906768, 33657023
2468	87614696 (4935, 4936)		Contains protein domain (PF00400) - kinase WD domain, G-beta repeat	kinase	29331824, 52644045, 265008, 264910, 265019, 21906765, 21906769, 265021
2469	86294397 (4937, 4938)				264288, 264628
2470	80223831 (4939, 4940)	80223831 (4939, 4940) Novel Protein sim. GBank gil5420389 emb CAB46680.1 - (4928-4938) (AJ243460) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	264592, 264692, 264555, 264556, 264557, 264558, 264559, 18108385, 264482
2471	91013681 (4941, 4942)	91013681 (4941, 4942) Novel Protein sim. GBank gil5419882 emb CAB46424.1 - (AL096749) DKFZp434G153 [Homo sapiens]		UNCLASSIFIED	65274572, 35696286, 29331827, 265007, 264592, 33109954, 265018, 265019, 264288, 55811957, 265020, 264693, 55811576, 56182323
		95060811 (4943, 4944) Novel Protein sim. GBank gil4929747[gb AAD34134.1 AF15189 - (AF151897) CGI-139 protein [Homo sapiens]		UNCLASSIFIED	264092, 60432049, 29331825, 60433356, 265010, 265011, 18108351, 264764, 264288, 264692, 65274620, 18108370, 18108372, 18108374, 264634, 18108385
2473	95421509 (4945, 4946)	95421509 (4945, 4946) Novel Protein sim. GBank gil4539009 emb CAB39630.1 - (AL049481) putative protein [Arabidopsis thaliana]			60424179, 65274572, 22278999, 60424269, 29331826, 265008, 60433356, 60433438, 265010, 18108351, 264448, 264288, 264687, 264689, 265021, 264692, 65274620, 60431528, 65274791, 264556, 56182323, 60432113
2474	94315616 (4947, 4948)	2474 94315616 (4947, 4948) Novel Protein sim. GBank gil3252827 (AC004382) - Unknown gene product [Homo sapiens]			65274572, 56994075, 264259, 29331826, 60170831, 265017, 265018, 265019, 264683, 264369, 265020, 264693, 264563, 264564

264259, 29331822, 265006, 265007, 265010, 265011, 264448, 264288, 264369, 264685, 264686, 18108357, 264768, 18108362, 264693, 18108370, 18108374, 18108383, 18108385, 264565, 264567, 264565, 264567	264259, 60424269, 66714117, 264905, 265006, 265008, 265008, 265009, 264758, 265010, 265011, 18108351, 264681, 264369, 264288, 264689, 21906767, 265020, 18108374, 264639, 18108382, 83373044, 18108385, 87168518	263978	265017	56994075, 22278999, 21906754, 264682. 21906765	264905, 264907, 264765	264269, 29331822, 29331824, 667144177, 29331827, 35696052, 264508, 264905, 264906, 264907, 2659018, 264905, 264906, 264907, 265008, 264906, 264906, 264907, 265008, 2649010, 33657402, 265018, 264760, 264760, 265011, 265017, 265018, 264760, 264762, 18108351, 264764, 264288, 264760, 264762, 18108351, 264764, 264288, 264760, 264686, 264693, 264629, 264693, 264636, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264637, 264638, 264907, 264908, 264907, 264908, 264907, 264908, 264408, 264288, 263967, 181083370, 18108385, 18108388, 264468, 264563	22278995, 22278996, 22278997, 22278999, 264259, 60432049, 29331824, 60432289, 29331827, 265007, 264910, 264593, 264600, 264603, 264604, 265019, 264448, 264288, 264685, 264686, 264693, 264689, 35695917, 265022, 264692, 264693, 56182323
J61 -	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	tm7	UNCLASSIFIED	UNCLASSIFIED UNCLASSIFIED	UNCLASSIFIED
1216486 (U48852) - HT protein Contains protein domain (PF00008) - tgf EGF-like domain							
	94315618 (4951, 4952) Novel Protein sim. GBank gil3252827 (AC004382) - Unknown gene product [Homo sapiens]			94314569 (4957, 4958) Novel Protein sim. GBank gi 1644232 db BAA11082 - (D67066) N-WASP Bos taurus]		#7393165 (4963, 4964) Novel Protein sim. GBank gij321249 pir S28407 - guanine nucleotide-exchange activator CDC25 homolog - mouse mucleotide - mouse (4965, 4966) Novel Protein sim. GBank gij321249 pir S28407 - guanine nucleotide - mouse (4965, 4966) Novel Protein sim. GBank gij321249 pir S28407 - guanine g4187774 (4967, 4968) Novel Protein sim. GBank gij321249 pir S28831 sp P39188 ALU1_HUMAN - IIII ALU SUBFAMILY	J VVARMING EN I RT !!!! 87786556 (4969, 4970) Novel Protein sim. GBank gil1185397 (U25281) - SH3 domain binding protein [Rattus norvegicus]
2475 94321693 (4949, 4950) Novel Protein sim. GBank gil [Cricetulus griseus]	2476 94315618 (4951, 4952)	177 20718974 (4953, 4954)	(2478 17659165 (4955, 4956)		2480 95295605 (4959, 4960)	87393165 (4963, 4964) 87731583 (4965, 4966) 94187774 (4967, 4968)	2485 87786556 (4969, 4970)

265017, 264555	22278996, 35696286, 22278997, 22278998, 22278999, 264092, 264259, 28331822, 35696052, 264108, 264905, 284907, 22331830, 264309, 264905, 284907, 29331830, 264909, 265006, 264511, 265008, 265009, 60433438, 21906754, 33109954, 87168559, 265018, 26481, 264288, 264687, 21906768, 21906768, 21906768, 21906768, 21906769, 3569502, 264534, 33657023, 264692, 33657109, 263972, 18108377, 35696423, 35695855, 60170334, 18108385, 56526486, 22279000, 22279002, 264563, 264482, 264565, 20281189, 18108391	264910, 264448, 264288, 264684, 264691, 264634	264686, 264693, 55811576, 22279002	264907, 265008, 22279002	264259, 29331826, 265008, 264762, 18108370, 18108376, 18108379	264488, 22278998, 22278999, 29331828, 264591, 33109954, 265017, 55811150, 21906764, 21906768, 264692, 60431528, 87168518, 60432113, 22279000
ļ	glycoprotein	UNCLASSIFIED	complementrecept	UNCLASSIFIED	lm7	transcriptfactor
	Ras family RAB-14 Ras family Ras family		Contains protein domain (PF00084) - complementrecept Sushi domain (SCR repeat)			
2486 87748978 (4971, 4972) Novel Protein sim. GBank gi 2662167 db BAA23715 - (AB007903) KIAA0443 Homo sapiens	95343105 (4973, 4974) Novel Protein sim. GBank gil464559 sp P35287 RB14_RAT RAS-RELATED PROTEIN RAB-14		82990585 (4977, 4978) Novel Protein sim. GBank gil4886439 emb CAB43355.1 - (AL050253) hypothetical protein [Homo sapiens]	2490 88069609 (4979, 4980) Novel Protein sim. GBank gi[258624 (AC003083) - Rap2 interacting protein-like; similar to U73941 (PID:g1916018) [Homo sapiens]	91242116 (4981, 4982) Novel Protein sim. GBank gij728832 sp P39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!	95308202 (4983, 4984) Novel Protein sim. GBank gil3355303 (AF001549) - Unknown gene product [Homo sapiens]
87748978 (4971, 4972)	95343105 (4973, 4974)	2488 87652451 (4975, 4976)	82990585 (4977, 4978)	88069609 (4979, 4980)		
2486	2487	2488	2489	2490	2491	2492

2483	95422415 (4985, 4986)	2483 95422415 (4985, 4986) Novel Protein sim. GBank gil4240307 db 18474932.1	Contains protein domain (PF01424) - struct	struct	18108394, 264887, 65274572, 56182575,
		(AB020716) KIAA0909 protein [Homo sapiens]	R3H domain		22278995, 56994075, 60432049, 29331822,
					29331824, 29331825, 29331826, 29331827,
					29146498, 264508, 264905, 264509, 264906,
					264907, 29331830, 264909, 264510, 265006,
					264511, 265007, 264512, 265008, 265009.
					264910, 21906754, 265011, 264600, 265017,
					265018, 264604, 264605, 265019, 55811150,
					264762, 18108351, 264681, 264448, 264683,
					264369, 264288, 18108355, 18108357,
_					264687, 21906765, 21906766, 21906767.
				-	21906768, 21906769, 265020, 264691,
					264692, 33657023, 33657349, 18108370,
					18108374, 18108376, 55810764, 18108379,
	_				65274791, 264630, 264632, 264634, 264635,
					264636, 264555, 264637, 264557, 264558,
					264639, 264559, 83373044, 18108385,
					87168518, 60432113, 22279000, 22279002,
					264482, 264566, 264486
2494				UNCLASSIFIED	264907, 264601
2495		94234551 (4989, 4990) Novel Protein sim. GBank gil5420389 emb CAB46680.1 -		collagen	263994, 22278997, 35696052, 264509,
		(AJ243460) proteophosphoglycan [Leishmania major]			264905, 264906, 264907, 264908, 264909,
					265006, 265009, 264595, 264604, 264448,
					264682, 264764, 264288, 264685, 264766,
					264769, 264689, 265020, 264692, 65274620,
					264629, 55810764, 35696423, 55811576,
_					264636, 264637, 18108385, 22279000,
					264564, 264567, 264486
2496		80018765 (4991, 4992) Novel Protein sim. GBank gil4808220 emb CAB42832.11 -		struct	29147620, 264905, 265006, 265007,
		(AL022315) dJ117715.1 (PUTATIVE novel protein) [Homo			18108348, 18108362, 18108370, 18108374,
		sapiens]			264555, 264556, 18108381, 18108383,
	7				18108388
2497	91723554 (4993, 4994)			UNCLASSIFIED	52644507, 22278996, 22278999, 29331824,
					29331828, 33657402, 21906754, 87168474,
					265019, 264369, 264689, 21906765,
					21906766, 21906767, 21906768, 265020,
					33657023, 18108376, 18108387
2498		87724633 (4995, 4996) Novel Protein sim. GBank gi 1200503 (U47924) - B [Homo sapens]		UNCLASSIFIED	29331827, 264512, 264910, 264288, 18108374, 35695855
2499	•	94685125 (4997, 4998) Novel Protein sim. GBank gi[3510234 (AC005581) -	Contains protein domain (PF00069) - kinase	kinase	264909, 55812038, 264631, 264637, 264558
		R31237_1, partial CDS [Homo sapiens]	Eukaryotic protein kinase domain		

WO 00/58473

PCT/US00/08621

(80)	2504 87868706 (5007, 5008) Novel Protein sim. GBank gij550420jemb CAA48220 - (X68101) trg [Rattus norvegicus]			264488, 52644507, 52645156, 52646842. 22278994, 264259, 52645080, 29331822. 29331824, 29331825, 29331826, 29331827. 35696052, 264906, 264908, 52644045, 265009, 60433356, 33657402, 60433438.
				265019, 264448, 264288, 264766, 52644229, 21906765, 21906766, 21906767, 21906768,
				52644150, 264692, 27486261, 27486262, 27486264, 27486264, 35695763, 35696423,
				35695855, 52644332, 56182323, 18108387, 87168518, 60432113, 22279002, 264564
\perp			UNCLASSIFIED	264605
Š Ž	2506 91232326 (5011, 5012) Novel Protein sim. GBank gil2137562 pir 149635 - mouse		nuclease	264488, 52644507, 52645156, 52646365, 66274672, 22278995, 66994075, 22278996
E	Dhm1 protein - mouse			22278997, 22278998, 22278999, 264259,
				60432049, 29331822, 29331825, 29331826,
				264593, 60433356, 55812038, 21906754,
				265011, 265017, 265018, 265019, 264605,
				264762, 18108351, 264448, 264288, 264768.
		-		21906765, 21906766, 21906767, 21906768, 21906769, 31606769, 316067617, 265020, 265021
				265022, 60170615, 33657023, 27486264,
				18108379, 35695855, 264637, 83373044.
				18108385, 87168518, 60432113, 22279000,
				264563, 264482, 264565
<u>ž</u>	1	Contains protein domain (PF00850) - histone	histone	264488, 263994, 264592, 264595, 264369,
<u>.</u>	gij5174489 ref NP_006035.1 pKIAA - histone deacetylase 6	Histone deacetylase family		264686, 264768, 33693917, 33692423, 264563
Ž	95315505 (5015, 5016) Novel Protein sim. GBank gil4826433 emb CAB42889.1 -		UNCLASSIFIED	22278995, 22278999, 60432049, 264259.
<u> </u>	(AL031447) dJ126A5.2.1 (novel protein) (isoform 1) [Homo			29331828, 265006, 265007, 60433438. 33657084, 265010, 265017, 265018, 265019,
<u> </u>	sapiens)			18108351, 264448, 18108354, 264369,
				18108359, 21906765, 21906769, 55811957,
			•	265020, 265022, 27486261, 33657349,
				18108377, 35695855, 60432113, 22279002, 264563, 264563
+3	Ovel Protein sim GBank gill 263289 (147856) - fibroin-4		UNCLASSIFIED	265007, 265008, 18108357, 264556, 264567
==	6/615/41 (3017, 3010) Novel Frotein Sint. Gbairk gij (203203 (047030) - iidoni 4 [Araneus diadematus]			
1				

65274572, 22278994, 22278996, 22278998, 22278999, 60432049, 264259, 29331822, 29331824, 29331825, 66714117, 60432289, 29331826, 3355970, 264908, 66712502, 265007, 264910, 6043335, 60433438, 21906754, 81168474, 265017, 265018, 264448, 264288, 21906765, 21906768, 21906769, 25811957, 35695917, 255020, 256027, 264691, 33657182, 27486262, 33657349, 18108370, 35695855, 264555, 56182323, 83373044, 60432113, 22279002	265017, 21906764, 265020, 264692	22278999, 264259, 29331825, 29331826, 29146499, 264907, 264909, 265006, 265008, 264591, 60432229, 21906754, 264763, 264683, 264666, 18108377, 264689, 264669, 18108370, 263972, 18108374, 264586, 22279000	60424179, 52645156, 18108394, 22278994, 35696266, 56994075, 22278996, 29331822, 29331824, 60424269, 29331825, 29331827, 33656970, 60431735, 33657084, 87168559, 265077, 264448, 264389, 56181562, 21906766, 21906769, 21906768, 21906769, 255021, 33657023, 18108366, 33657109, 27486261, 27486262, 33657349, 18108374, 55810764, 35698423, 56182323, 264558, 18108385	264510		264259, 29146498, 264905, 264288, 29148629, 35695917, 27486261, 264634	264091, 29331824, 29331825, 29331826, 29331828, 35696052, 264508, 264509, 264909, 264905, 264900, 264907, 264907, 264909, 264511, 264910, 33657402, 264757, 33109954, 265017, 265018, 264768, 264769, 264763, 264763, 26468, 55811576, 33657109, 33657182, 264628, 55811576, 35696423, 264631, 264634, 264637, 264638, 2279002, 264564
dna_ma_bind		Iransport	UNCLASSIFIED	UNCLASSIFIED	. tm 7	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF00097) - transport Zinc finger, C3HC4 type (RING finger)	·		Contains protein domain (PF00001) - tm7 7 transmembrane receptor (rhodopsin family)		
2510 95421379 (5019, 5020) Novel Protein sim. GBank gil3293537 gb AAC25762.1 - (AF071059) zinc finger RNA binding protein [Mus musculus]	87384281 (5021, 5022) Novel Protein sim. GBank gil4323152 gb AAD16228.1 - (AF096863) Ets-protein Spi-C [Mus musculus]		95357843 (5025, 5026) Novel Protein sim. GBank gil3004657 (AF01777) - bobby sox [Drosophiia melanogaster]	88094578 (5027, 5028) Novel Protein sim. GBank gij2258437 (AF008197) - syncollin [Rattus norvegicus]	87994509 (5029, 5030) Novel Protein sim. GBank gil3757727 emb[CAA18783 - (AL022727) dJ80119.7 (olfactory receptor-like protein (ths6M1-3)) (Homo sapiens)	(2)	87784966 (5033, 5034) Novel Protein sim. GBank gil4220527 emb CAA23000 - (AL035356) putative protein [Arabidopsis thaliana]
510 95421379 (5019, 502			2513 95357843 (5025, 5020	2514 88094578 (5027, 5020	2515 87994509 (5029, 5030	2516 87786908 (5031, 5032)	2517 87784966 (5033, 503

	264259, 66714117, 29331826, 29331827, 29331828, 264907, 66712502, 265006, 265008, 264594, 265010, 265011, 265018, 264288, 21906769, 265020, 60431528, 55811576, 65274791, 264632, 264555, 264636, 22279002, 264564	264259, 264563 265020, 264563	224488, 264489, 263994, 652/45767, 22278995, 22278998, 264259, 29331822, 29331826, 264508, 264509, 264509, 264906, 264907, 66712502, 264511, 265006, 265007, 264591, 264592, 264593, 264594, 264596, 264764, 264684, 26448, 264763, 264682, 264686, 21906768, 58811957, 264682, 264693, 27486261, 18108370, 264628, 264629, 18108374, 55811576, 35696423, 35695855, 264632, 264558, 18108385, 65274727, 60432113, 264563, 264564,	264489, 22278997, 20281171, 21906/54, 35695917, 263967, 263976, 263981, 20281169 263969	56994075, 22278996, 35696286, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 29331826, 29331828, 29331830, 56182435, 264512, 265008, 60170831, 33657402, 265010, 87168559, 265019, 264288, 21906765, 21906769, 35695917, 265020, 265021, 265022, 52644150, 264681, 33657023, 33657109, 27486281, 35696423, 65274791, 264559, 83373044, 56526486, 87168518, 264567
UNCLASSIF	kinase	transport	histone	transport	
Contains protein domain (PF00018) - UNCLASSIFIED SH3 domain			Contains protein domain (PF00850) - histone Histone deacetylase family		Contains protein domain (PF01753) - MYND finger
Novel Protein sim. GBank gil4929591[gb AAD34056.1 AF15181 - (AF151819) CGI-61 protein [Homo sapiens]	2519 94326180 (5037, 5038) Novel Protein sim. GBank gil4263748[gb]AAD15420 - (AC004883) similar to KIAA0766; similar to PID:g3882253 [Homo sapiens]	Novel Protein sim. GBank gij4826722 ref NP_005085.1 pFATP - fatty acid transport protein 4	95316244 (5041, 5042) Novel Protein sim. GBank gi 5174489 ref NP_006035.1 pKIAA - histone deacetylase 6	87754052 (5043, 5044) Novel Protein sim. GBank gil4580011[gb]AAD24201.1[U81002 (U81002) TRAF4 associated factor 1 [Homo sapiens]	95340469 (5047, 5048) Novel Protein sim. GBank gi[1809327 (U76374) - skm- 95340469 (5047, 5048) BOP2 [Mus musculus]
2518 94147410 (5035, 5036) Novel Protein sim. GBank gil4929591gb AAD34056. protein [Homo sapiens]	94326180 (5037, 5038)	2520 87413235 (5039, 5040) Novel Protein sim. GBank gil4826722 ref NP_005086 protein 4	95316244 (5041, 5042)	87754052 (5043, 5044)	95340469 (5047, 5048)
2518	2519	2520	2521	2522	2524

					0.000.00 000000000000000000000000000000
2525	94126928 (5049, 5050)	otic		Kinase Kinase	264488, 7227 6991, 7227 8999, 60432049, 60432289, 2043289, 2043289, 2043289, 2043289, 24369, 255008, 2561208, 21906756, 255007, 21906769, 23695917, 265020, 265021, 33657109, 60431528, 83373044, 60432113, 22279000, 22279002, 264565
7526		4589628 dbj BAA76836.1 -	Contains protein domain (PF00238) - Iribosomalprot	ribosomatprot	60424179, 264768, 264687, 264769, 264689, 6574572, 21906767, 5618275, 21906768, 21906768, 21806768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906268, 21906268, 2190626, 2190622, 264592, 264591, 60432049, 264259, 264097, 2365702, 264591, 60432049, 264259, 246097, 2365702, 264509, 264905, 264907, 218108372, 18108374, 56182435, 18108372, 18108374, 56182435, 264909, 18108372, 18108374, 56182435, 265009, 264097, 264515, 265008, 265009, 264097, 264515, 265008, 264595, 264592, 60431735, 264598, 264592, 55112038, 264596, 264598, 264588,
2527		88094580 (5053, 5054) Novel Protein sim. GBank gi[2258437 (AF008197) - syncollin [Rattus norvegicus]		UNCLASSIFIED	264510, 264512, 264630, 264591, 264592, 264259, 264259, 264259, 264503, 264605, 18108351, 264565, 264369, 18108354
2528		88078380 (5055, 5056) Novel Protein sim. GBank gil2085786 (AC002086) - similar to zinc finger 5 protein from Gallus gallus, U51640 (PID:91399185) [Homo sapiens]	Contains protein domain (PF00651) - dna_ma_bind BTB/POZ domain	dna_rna_bind	55812038, 265017, 264689, 35695917, 35695763, 60431528, 60432113, 22279002
2529		86670926 (5057, 5058) Novel Protein sim. GBank gil3786433 (AF098505) - similar to Arabidopsis thaliana male sterility protein 2 (SW:Q08891) [Caenorhabditis elegans]		synthase	264908, 264769, 265020, 265021, 18108383
2530	80259978 (5059, 5060)	-			264369, 264556
2531				UNCLASSIFIED	29331822, 29331824, 60432289, 264508, 264509, 264906, 265011, 264769, 21906768, 33657023, 87168518, 22279000
2532		87419778 (5063, 5064) Novel Protein sim. GBank gil2864625 emb CAA16972 - (AL021811) putative protein [Arabidopsis thaliana]			264593
2533		87000255 (5065, 5066) Novel Protein sim. GBank gil437181 (U02289) - GTPase- activating protein [Caenorhabditis elegans]		UNCLASSIFIED	264555

22279000 2227900 22279000 22279000 222790	335355 (3007, 3008	2534 87332322 (5067, 5068) Novel Protein sim. GBank gi[3452473 (AF084205) -	UNCLASSIFIED	264259, 35696052, 264905, 265017,
HUMAN - III! ALU SUBFAMILY Winase		serine/threonine protein kinase TAO1 [Rattus norvegicus]	-	21906769, 265020, 265022, 33657109,
PUTATIVE novel protein) HUMAN - IIII ALU SUBFAMILY IPHERC - guanine nucleotide UNCLASSIFIED GION	(5069, 5070	Novel Protein sim. GBank gil4468311 lemb CAB37992 -		65274572 35696286 60432289 29331828
HUMAN - IIII ALU SUBFAMILY [PHERC - guanine nucleotide UncLASSIFIED UNCLASSI		(AL031432) dJ465N24.2.1 (PUTATIVE novel protein)		66712502, 265006, 60432229, 265017.
HUMAN - III! ALU SUBFAMILY IPHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION UNCLASSIFIED		(isoform 1) [Homo sapiens]		265018, 265019, 264288, 264369, 264689,
HUMAN - IIII ALU SUBFAMILY IPHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION UNCLASSIFIED				21906768, 265020, 265021, 264636,
HUMAN - III! ALU SUBFAMILY IPHERC - guanine nucleotide UNCLASSIFIED GION	7 15074 5072	No. of Control of Cont		60170394, 22279002
IPHERC - guanine nucleotide UNCLASSIFIED GON	0 (307 1, 3072		kinase	18108398, 56182575, 35696286, 22278997,
Interior of the control of the con		SO MAN DAILED THE SALES AND THE ALC SUBTAMILY		22278999, 60432049, 264259, 29331824,
Intercondition Phercondition Pherconditi		OF WARMING ENIRY !!!		29331826, 29331827, 29331828, 264905,
IPHERC - guanine nucleotide UNCLASSIFIED UNCL				264511, 265009, 264910, 264596, 52646317,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				18108351, 264681, 264683, 18108354,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				264288, 264687, 264769, 264689, 21906765,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GOON				21906766, 21906767, 265021, 52645129,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				33657109, 18108374, 18108380, 56182323,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				18108381, 18108388, 87168518, 60432113,
pHERC - guanine nucleotide UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				22279000, 22279002, 264567, 18108391
IPHERC - guanine nucleotide	3 (5073, 5074,		ubiquitin	65274572, 35696286, 29331822, 29331825,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UI fly (Drosophila WOUSE - SER/THR-RICH GION		_		29331827, 29331828, 35696052, 264906,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UI fly (Drosophila WOUSE - SER/THR-RICH GION		exchange factor p532		66712502, 264909, 265008, 265011, 264760,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				264288, 264685, 35695917, 60170615,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION				264691, 33657023, 65274620, 33657109.
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UI fly (Drosophila UI fly (Drosophila WOUSE - SER/THR-RICH GION		-		18108374, 35696423, 35695855, 264636,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UI fly (Drosophila UI fly (Drosophila UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED GION	4 15075 5070			264558, 60170394, 56182323, 83373044
UNCLASSIFIED UNCLASSIFIED UNI fly (Drosophila WOUSE - SER/THR-RICH GION	4 (30/3, 30/6		UNCLASSIFIED	
1362647 pir S53876 - sex- uit fly (Drosophila ui Ovicial Sified UNCLASSIFIED GION	6 (5077, 5078,		UNCLASSIFIED	22278996, 22278999, 29331822, 29331825,
1362647 pir S53876 - sex- uit fly (Drosophila uit dy (Drosophila UNCLASSIFIED GION				29331828, 29146499, 264908, 264112,
UNCLASSIFIED uit fly (Drosophila uit fly (Drosophila UNCLASSIFIED GION				60170831, 87168559, 264604, 265019,
UNCLASSIFIED uit fly (Drosophila uit fly (Drosophila UNCLASSIFIED GION				264685, 264766, 87168518, 22279000,
UNCLASSIFIED uit fly (Drosophila uit construction of the constru	0000			264565, 264566
MOUSE - SERTHR-RICH	5 (5078, 5080)		UNCLASSIFIED	22278997, 29331828, 265008, 265009,
MOUSE - SERTHR-RICH		regulated protein janus A - full fly (Drosophila		264758, 265010, 18108351, 264683, 264288,
UNCLASSIFIED LEGION		bseudodoscura		21906765, 35695917, 265020, 18108374,
MOUSE - SER/THR-RICH LEGION	(5081 5082)	Novel Protein sim CBank		264567
	14000 (1000)	_	UNCLASSIFIED	264486, 264768, 264689, 264511, 202811/1, 264634, 264635, 264604, 264630, 26234834
-		PROTEIN TAO IN DOCK BEGION		204034, 204035, 204091, 204039, 29331824,
			-	264603, 264604, 264905, 264907, 264908,

2542	95298162 (5083, 5084	95298162 (5083, 5084) Novel Protein sm. GBank gil5225320[gblAAD40850.1]AF08310 - (AF083107) sirtuin type 2 [Homo sapiens]	Contains protein domain (PF00220) - UNCLASSIFIED Neurohypophysial hormones, N-terminal Domain	UNCLASSIFIED	26448, 18108394, 52646365, 52646842, 65274572, 22278894, 36996286, 22278996, 264259, 52645080, 29331822, 29331824, 29331827, 29331827, 29331827, 356900, 264900, 264900, 264900, 264900, 264900, 265001, 265001, 265001, 265018, 265009, 266910, 265011, 265018, 265019, 18108351, 264683, 264288, 264687, 5264637, 25644150, 21906766, 21906767, 21906769, 33657109, 52645129, 33657182, 27488261, 27486243, 33657349, 33695763, 18108374, 35696423, 3569585, 264631, 264634, 264635, 264558, 83373044, 264634, 264658, 87168518, 264563, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 26456, 2645636, 264563, 264564, 26456, 26
		94139086 (5085, 5086) Novel Protein sim. GBank gil5419857 emb CAB46374.1 - (AL096723) hypothetical protein [Homo sapiens]	Contains protein domain (PF00076) - RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)		65274572, 56182575, 22278999, 264259, 29331826, 264907, 264510, 264511, 264592, 264569, 264569, 264569, 264569, 264569, 264684, 264764, 264369, 21906767, 21906768, 60170615, 264692, 264693, 55811576, 65274791, 264639, 18108385, 60432113, 22279000
2544	94218549 (5087, 5088	94218549 (5087, 5088) Novel Protein sim. GBank gi[2498110]sp]G631911AEGP_RAT - APICAL ENDOSOMAL MAM domain. GLYCOPROTEIN PRECURSOR	Contains protein domain (PF00629) - glycoprotein MAM domain.	glycoprotein	18108397, 52646365, 22278997, 264259, 60432049, 29331822, 29331825, 29331826, 29331827, 29331826, 29331827, 29331828, 265408, 265006, 265007, 265008, 87168559, 265017, 265018, 265619, 18108351, 264484, 264686, 265021, 18108370, 18108374, 18108376, 18108381, 18108385, 26526486, 22279000, 264482, 264563, 264567
745	87742645 (5089, 5090	2545 87742645 (5089, 5090) Novel Protein sim. GBank gi]3327046 dbj BAA31591 - (AB014516) KIAA0616 protein [Homo sapiens]			29331825, 264906, 265009, 60170831, 265017, 264369, 21906767, 60170615, 264692, 33657109
2546	88093861 (5091, 5092	88093861 (5091, 5092) Novel Protein sim. GBank gi 2996032 (AF054586) - brain finger protein [Rattus norvegicus]	Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)	UNCLASSIFIED	29331824, 265007, 22279002

2547	94143869 (5093, 5094	2547 94143869 (5093, 5094) Novel Protein sim. GBank gil4929607[gblpAD34064.1 AF15182 - (AF151827) CGI-69 protein [Homo sapiens]	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	ransport	264488, 18108394, 52646842, 18108397, 56182575, 22278995, 56984075, 22278996, 22278997, 22278999, 264259, 29331822, 29331824, 29331828, 264259, 29331822, 29331824, 29331828, 264104, 264508, 2648045, 264809, 26482435, 265006, 264901, 264001, 265001, 265001, 265001, 264001, 265010, 265011, 264600, 264601, 265018, 265019, 264760, 18108354, 264682, 26448, 265016, 265016, 265011, 264600, 264601, 265018, 264280, 264389, 21906765, 21906765, 21906765, 21906769, 265021, 264691, 18108374, 265022, 264690, 264691, 18108374, 265022, 264690, 264691, 18108374, 264693, 27486261, 18108370, 18108362, 264693, 264659, 264659, 264659, 264659, 264659, 264559, 264659, 264559, 264559, 264559, 264559, 264559, 264559, 264559, 264559, 264569, 264566, 264666, 264566, 264666, 264666, 264666, 264666, 264666,
2548	88179079 (5085, 5096) ,			UNCLASSIFIED	264488, 18108394, 52646365, 22278994, 35696286, 56994075, 22278997, 22278999, 264259, 29331822, 29331824, 29331825, 29331826, 29331826, 29331827, 29331828, 29331827, 29331828, 26182435, 264511, 265007, 264512, 60433356, 87168559, 264684, 264369, 52644229, 265021, 33657023, 264692, 18108384, 18108385, 60432113, 22279000, 22279002, 264563, 264567
2549		94196893 (5097, 5098) Novel Protein sim. GBank gij728837 sp P39194 ALU7_HUMAN - III! ALU SUBFAMILY SQ WARNING ENTRY III!	Contains protein domain (PF00412) - struct LIM domain containing proteins	struct	56182575, 22278996, 22278997, 22278998, 22278999, 264259, 264508, 264908, 29331830, 265009, 265010, 265018, 264688, 21906764, 21906765, 21906766, 21906767, 21906769, 265021, 52644150, 264691, 18108368, 60431602, 18108376, 35696423, 56182323, 18108387, 264567
2550	87778584 (5099, 5100	87778584 (5099, 5100) Novel Protein sim. GBank gi 2143886 pir 152523 - nucleoporin p62 homolog - rat (fragment)		UNCLASSIFIED	56182575, 29331822, 264105, 264512, 18108351, 35695917, 264637, 264638

	95308400 (5101, 5102)	[2551 195308400 (5101, 5102) Novel Protein sim. GBank gil4337103 gb AAD18079 -	Contains protein domain (PF00561) - UNCLASSIFIED	UNCLASSIFIED	18108396, 65274572, 22278995, 22278998,
		(AF129756) NG26 [Homo sapiens]	alpha/beta hydrolase fold		22278999, 264259, 29331822, 29331824,
					29331825, 60432289, 29331826, 29331827,
					264905, 56182435, 265007, 60433438,
					55812038, 21906754, 65274444, 265017,
		~~~			265018, 264605, 265019, 264288, 21906766,
					21906768, 21906769, 265020, 60170615,
					264693, 33657109, 35696423, 264638,
					56182323, 83373044, 22279000
	95332620 (5103, 5104)			UNCLASSIFIED	56182575, 35696286, 29331824, 29331826,
					35696052, 29331828, 264508, 264907,
			,	T.	56182435, 265008, 264591, 33109954,
					264760, 55811957, 35695917, 33657023.
					33657109, 18108374, 55811576, 35696423,
					35695855, 56182323, 264558
	95308243 (5105, 5106)	95308243 (5105, 5106) Novel Protein sim. GBank		UNCLASSIFIED	264686, 264488, 263976, 264768, 29331826,
		gil771058jspjr34797j110_MOUSE - SERVIHK-KICH			33095032, 33690423, 204001, 204311, 364602, 364010, 364634, 364760, 364565
_					264762 264910, 204034, 204700, 204333, 264762 264906 264502 264691 264566
					264908, 264684, 264567, 264909, 264766
_	87761520 (5107, 5108)	87761520 (5107, 5108) Novel Protein sim. GBank		cadherin	22278997, 29331822, 264508, 21906769,
		gil728835 sp P39192 ALU5_HUMAN - IIII ALU SUBFAMILY			33657023, 33657109, 56182323
2566	07637664 (6400 6440)	SC WANKING ENTRY !!!			00004004
	6/62/331 (3108, 3110)	o/oz/331 (3109, 3110) Novel Protein sim. GBank gij4884319jemb CAB43260.1j -  (AL050084) hypothetical protein [Homo sapiens]		nuclease	29331024, 203972
2556		87645533 (5111, 5112) Novel Protein sim. GBank gil4106984 (AC003038) -	Contains protein domain (PF00514) -	UNCLASSIFIED	22278998, 264509, 33657402, 264683,
	į	R30923_1 [Homo sapiens]	Armadillo/beta-catenin-like repeats		264684, 264766, 264689, 33657023, 33657109, 35695855, 264558, 264567
	78437803 (5113, 5114)				264595
2558	87617591 (5115, 5116)	87617591 (5115, 5116) Novel Protein sim. GBank gij119110jspjP03211jEBN1_EBV - EBNA-1 NUCLEAR PROTEIN		UNCLASSIFIED	22278997, 29331824, 66714117, 29331825, 264906, 264511, 265018, 264448
2559		88095382 (5117, 5118) Novel Protein sim. GBank gil4538998 emb CAB39619.1		UNCLASSIFIED	22278997, 29331822, 29331828, 60433356,
		(AL049481) AIG1-like protein [Arabidopsis thaliana]			265011, 264288, 264765, 264766, 264769, 21906765, 21906766, 60432113, 264482
2560		87994530 (5119, 5120) Novel Protein sim. GBank gil5051399 emb CAB44995.1  - (AL078630) 573K1.3 (mm17M1-4 (novel 7 transmembrane receptor (rhodopsin family) (olfactory receptor LIKE) protein)) [Mus musculus]	Contains protein domain (PF00001) - 7 transmembrane receptor (rhodopsin family)		-
	88176575 (5121, 5122)	88176575 (5121, 5122) Novel Protein sim. GBank gij5326825jgbJAAD42056.1JAF04495 - (AF044953)		UNCLASSIFIED	22278995, 35696286, 22278996, 22278997, 22278998, 22278999, 264259, 29331822,
		NADH:ubiquinone oxidoreductase PGIV subunit [Homo saniens]			29331824, 29331826, 29331827, 29331828, 265007, 60432229, 87168559, 265017,
					265018, 265019, 264689, 21906766,
_					21906769, 35695917, 265020, 33657023,
					33657109, 18108374, 264634, 264559,
_					18108385, 87168518, 22279002

2562 87645539 (5123, 5124) Novel Protein sim. GBank gi	Novel Protein sim. GBank g	14106984 (AC003038) -		UNCLASSIFIED	56994075, 22278996, 22278997, 22278999,
K30923_1 [Tomo Sapiens]	K30323_1 (Homo sapiens)				264239, 29331622, 60432289, 33657402, 60433356, 21906765, 55811957, 60170615, 33657023, 264693, 35695855, 87168518
88095497 (5125, 5126) Novel Protein sim. GBank gil4886447 emb CAB43371.1  -   (AL050270) hypothetical protein [Homo sapiens]	Novel Protein sim. GBank gil4886447   Protein   Novel Protein sim. GBank gil486447   Novel Protein   Novel Pro	371.1 -		UNCLASSIFIED	264488, 35696286, 22278999, 264259, 29331822, 29331824, 35696052, 264508,
					264907, 264908, 264909, 52644045, 264510, 264511, 265009, 264910, 264591, 264593,
					33657402, 265017, 265018, 265019. 18108351, 264686, 21906767, 21906768,
					55811957, 35695917, 265020, 264691,
					264693, 27486262, 264628, 18108374,
			-		35696423, 35695855, 264632, 264634,
			-		264635, 264639, 264558, 18108384,
					87168518, 22279000, 22279002, 264482,
00503703 /5437 £430\ N=1==== O=====					264563, 264565, 264566, 264486
2004 Jougus/ 63 (3127, 3126) NOVEI Protein Sim. GBank				sulfotransferase	29331822, 265007, 265010, 265019, 264769.
gij1352944 sp P47179 YJ9P_YEAST - HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR	gil 352944 sp P47179 YJ9P_YEAST - HYPO 118.4 KD PROTEIN IN BAT2-DAL5 INTERGE PRECURSOR	THETICAL ENIC REGION			55811576, 56182323
85530906 (5129, 5130)				UNCLASSIFIED	66714117, 264909, 263978, 264632
80224956 (5131, 5132) Novel Protein sim. GBank gil628012 pir  A5 mvr 4 - rai	Novel Protein sim. GBank gij628012 pir  A5 myr 4 - rat	3933 - myosin l	628012 pir  A53933 - myosin 1   Contains protein domain (PF00063) - struct   Magein bead (motor domain)	struct	18108370, 35695855, 264556, 264558,
2567 86143590 (5133, 5134)	-				265020, 60170615
91233099 (5135, 5136)	Novel Protein sim. GBank			synthase	60424179, 18108394, 56181686, 56994075,
4,	4,	CAEEL - HYPOTHETICAL 33.8			22278999, 264490, 264259, 29331822.
KD PROTEIN R10E11.4 IN CHROMOSOME II		==			56182181, 29331824, 60424269, 29331825,
					29331826, 29331828, 264509, 29331830,
					265007, 265008, 265009, 33657402, 265010,
				•	265011, 265019, 264448, 264683, 264288,
					18108354, 264769, 21906766, 21906767,
				•	35695917, 265021, 33657023, 18108362,
					33657109, 33657182, 35695763, 60431528.
					55810764, 18108379, 83373044, 18108385,   60432113, 264482
					00432113, 204402

18108394, 56182575, 56181686, 22278995, 22278996, 56994075, 35696286, 22278997, 22278998, 22278999, 264259, 60432049, 29331822, 5618181, 29331824, 29331825, 60432289, 264269, 60432289, 265087, 264905, 264910, 60432229, 264592, 60433356, 60433438, 21906764, 87168559, 265017, 265018, 265019, 264682, 264448, 265017, 265018, 265019, 264682, 264448, 265021, 265019, 264690, 264691, 265021, 265022, 52644150, 264690, 264691, 264692, 264693, 65274620, 263967, 3569563, 264693, 65274620, 263967, 3569563, 3695863, 3693676, 3589376, 3589376, 3589378, 3589385, 264558, 18108381, 56182323, 18108382, 22279000, 264566	22278996, 29331822, 29331824, 66714117, 29331825, 60432289, 29331827, 35696052, 264907, 265009, 264758, 33109954, 265019, 264686, 264689, 265020, 265021, 264691, 264692, 264693, 18108374, 35695855, 264634, 56182323, 264639, 60170394, 83373044, 22279002, 264482	264508, 264905, 264907, 264828, 18108351, 264555, 264556, 264557, 264558, 264559	35696286, 29331827, 35696052, 264100, 264104, 264110, 264592, 21906754, 29148627, 29148627, 263972, 263974, 18108374, 263976, 35695855, 60170394, 264559, 18108385
franscriptfactor	phosphatase	struct	UNCLASSIFIED
Contains protein domain (PF00170) - transcriptfactor	Contains protein domain (PF01240) - phosphatase Protein phosphatase 2A regulatory subunit PR55		
2569 95313764 (5137, 5138) Noval Protein sim. GBank gil2599560[gb]AAB84166.1  - (AF029674) basic leucine zipper protein LZIP [Homo sapiens]	2570 94136754 (5139, 5140) Novel Protein sim. GBank gil4758954[ref]NP_004567.1[pPPP2 - protein phosphalase 2 (formerly 2A), regulatory subunit B (PR 52), beta isoform	87733750 (5141, 5142) Novel Protein sim. GBank gil732218 sp p34609 YO60_CAEEL - HYPOTHETICAL 128,6 KD PROTEIN ZK1098.10 IN CHROMOSOME III	2572 8,7627560 (5143, 5144) Novel Protein sim. GBank gil4884319 emb CAB43260.1  - (AL050084) hypothetical protein [Homo sapiens]
95313764 (5137, 5138) [	1136754 (5139, 5140)	37733750 (5141, 5142)	87627560 (5143, 5144) i
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3	105040000 15445		(Occopan)		SESCOPOS CESTEDS CESTOCOCO COSTOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOCOC
5767	93313928 (3143, 3140)	23/3   83313828 (3149, 3149) NOVE Protein Sim. GBank	Contains protein domain (Produced) - complement	complement	504400, 00424119, 00214912, 30102919, 56181686 22278005 56004075 22278097
					60432049 264259 29334822 29331824
					29331825, 60432289, 29331826, 29331827,
					29331828, 264104, 264107, 264508, 264906,
					29331830, 264909, 264510, 265006, 264512,
					265008, 265009, 264910, 264591, 264592,
					60432229, 264593, 60433356, 264594,
					60433438, 264595, 55812038, 264759,
					21906754, 87168474, 265010, 265011,
					87168559, 265017, 265018, 265019, 264761,
					264762, 264763, 264764, 264369, 264288,
					264685, 264766, 264686, 264687, 264688,
					264769, 56181562, 264689, 21906765,
					21906766, 21906767, 29148627, 21906768,
					21906769, 265020, 265021, 265022,
					60170615, 264690, 52644150, 264691,
					264692, 33657023, 65274620, 18108365,
					18108368, 27486265, 60431602, 264629,
					60431528, 263976, 65274791, 35695855,
					20281071, 60431850, 264637, 264638,
_					264558, 264639, 56182323, 60170394,
					83373044, 18108384, 87168518, 60432113,
					264482, 264564, 264565, 264566, 264567
2574	-	94746814 (5147, 5148) Novel Protein sim. GBank gij3334982 (AC005306) -	Contains protein domain (PF00651) - UNCLASSIFIED	UNCLASSIFIED	22278995, 264259, 60432289, 29331827,
	_	R27216_1 [Homo sapiens]	BTB/POZ domain		29331828, 33656970, 264908, 265008,
					264910, 264591, 33657402, 265018, 265019,
					264448, 264764, 264369, 264288, 18108357,
					21906765, 21906766, 21906768, 55811957,
					60170615, 264691, 33657023, 264693.
					33657109, 33657182, 27486261, 27486264,
					33657349, 264636, 264555, 83373044.
					18108385, 264482
2575				UNCLASSIFIED	264910, 264601, 264683, 264689, 264080
		gil4929729 gb AAD34125.1 AF15188 - (AF151888) CGI-130	0		
	_	protein [Homo sapiens]	C	- (1)	254250 25505059 264006 50422439
2576		95357881 (5151, 5152) Novel Protein sim. GBank		npidnitin	Z64Z59, 33696U5Z, Z649U6, 6U433436.
		gi 4680681 gb AAD27730.1 AF13295 - (AF132955) CGI-21			264581, 18108351, 264288, 52644150.
		protein (Homo sapiens)	nydrotases ramily 2		204020, 33030423
2577	86996621 (5153, 5154)	86996621 (5153, 5154) Novel Protein sim. GBank gil4337103 gb AAD18079  -		UNCLASSIFIED	29331825, 265018, 265019, 264685
į	-	וער ובשוטטן ועסבט (וויטווט פמטופווט)			264480 264006 264008 2646040 264606
8/07	87780941 (5155, 5156)			UNCLASSIFIED	204400, 204900, 204900, 204910, 204990, 204603 264604 264606 264768 21006769
					264628, 284630, 264634, 264639, 264563
2579	87292879 (5157 5158)			UNCL ASSIFIED	29331822, 29331824, 264767
	_				

	88166788 (5159, 5160)	2560   88166788 (5159, 5160)   Novel Protein sim. GBank gi 2588628 (AC003080) - Similar to KIAA0299; 60% similarity to AB002297 (PID:g2224539)			265007, 265018, 264762
_		[Homo sapiens]			
8	7899048 (5161, 5162)	87899048 (5161, 5162) Novel Protein sim. GBank gi 4406642 gb AAD20049  -	Contains protein domain (PF00595) - collagen	collagen	56994075, 29331824, 29331826, 29331828,
		(AF131809) Unknown [Homo sapiens]	PDZ domain (Also known as DHR or GLGF).		264905, 60433356, 60433438, 264758, 87168559, 21906769, 265022, 35695855,
					263981
_	87786789 (5163, 5164)	87786789 (5163, 5164) Novel Protein sim. GBank gi 2739367 (AC002505) - putative		eph	264488, 264907, 264908, 264910, 264764.
		phosphatidylinositol-4-phosphate 5-kinase [Arabidopsis Ihaliana]			264684, 264766, 284638, 264555, 264565
2583	91220850 (5165, 5166)	91220850 (5165, 5166) Novel Protein sim. GBank gil4378112 emblCAA16521.1  -	Contains protein domain (PF00047) - transcriptfactor	transcriptfactor	56181686, 264259, 264510, 264512, 264591,
		(AL021578) dJ453C12.2 (similar to transcription factor RBP- Immunoglobulin domain	- Immunoglobulin domain		264592, 264593, 264594, 264595, 264596,
		L) [Homo sapiens]	,		264603, 264629, 55810764, 264630, 264637.
	80430041 (5167 5168)				2500000 250000 350000 350000 3500000
				UNCLASSIFIED	264908. 264910, 264768, 264693. 16106374, 55811576, 56182323
2585		80436126 (5169, 5170) Novel Protein sim. GBank gi 2736151 (AF021935) - mytonic		kinase	264768
		dystrophy kinase-related Cdc42-binding kinase [Rattus novegicus]			
2586	91226136 (5171, 5172)				22278998, 264259, 29331822, 29331824,
					29331827, 29331828, 264906, 265007,
					265009, 264591, 60433356, 33657402,
		-			265018, 264762, 264288, 21906766,
					21906767, 21906769, 265022, 264691,
					83373044, 56526486, 22279002
	80430943 (5173, 5174)				264908, 265019, 264768, 264693, 55811576,
	1000 JE 141 300 1000				56182323
	85515607 (5177, 5178)	2589   85515607 (5177, 5178)   Novel Protein sim. GBank qii3021598lemb CAA71415  -		UNCLASSIFIED	264564 35696052, 264905, 264906, 264907, 264908.
	•	(Y10389) nuclear protein [Xenopus laevis]			264909, 265009, 265018, 264769, 35696423.
_					264636
	87054526 (5179, 5180)	87054526 (5179, 5180) Novel Protein sim. GBank gi[2104689 (U92793) - alpha	Contains protein domain (PF01055) - glucoamylase	glucoamylase	22278995, 29331830, 265008, 265010,
_		glucosidase II, alpha subunit [Mus musculus]	Glycosyl hydrolases family 31		265017, 264639
_	94192167 (5181, 5182)	94192167 (5181, 5182) Novel Protein sim. GBank		ebh	264259, 29331822, 264106, 264906,
		gi 5702202 gb AAD47199.1 AF12916 - (AF129166) long-			56182435, 265007, 265008, 33109954,
		chain acyl-CoA synthetase 5 [Homo sapiens]			264448, 55811957, 265020, 18108370,
					55811576, 22279002

	1 00 11 11 11 11 11 11		transact	18108307 56182575 35506286 55004075
322040 (3103, 3	85552646 (5165, 5164) NOVEL FIGURIN SIM. GEBAIN			264259, 29331822, 29331824, 29331826,
	HEART PROTEIN			60432289, 29331827, 29331828, 264906,
				264909, 265007, 265008, 264910, 60432229,
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				264766, 265021, 60170615, 33657023,
				33657109, 264629, 35696423, 35695855,
				264557, 264638, 60170394, 56182323,
				83373044, 56526486, 87168518, 264563,
				264482, 264565
7754416 (5185,	87754416 (5185, 5186) Novel Protein sim. GBank	1)	tm7	22278999, 29331825, 264758, 21906754,
	gil4929729lgbJAAD34125.1JAF15188 - (AF151888) CGI-130	-130		52646317, 265010, 18108351, 264288,
	protein [Homo sapiens]			264369, 21906768, 264693, 18108370,
				264637, 264638, 264482
95305758 (5187,			UNCLASSIFIED	264488, 18108398, 56182575, 35696286,
	gij4929587lgbjAAD34054.1 AF15181 - (AF151817) CGI-59	-26	-	22278997, 264093, 264259, 29331822,
	protein [Homo sapiens]			29331825, 66714117, 29331826, 264905,
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				87168474, 265017, 264760, 264448, 264764,
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79561676 (5189, 5190)	5190)		UNCLASSIFIED	264692
37538637 (5191,	87538637 (5191, 5192) Novel Protein sim. GBank gil4309681 gb AAD15478  -		UNCLASSIFIED	22278999, 264259, 265018, 264448, 265021,
	(AC006930) R33423_1 [Homo sapiens]			60431528
94784089 (5193, 5194)	⊢	7	UNCLASSIFIED	264905, 264509, 264908, 264762, 264766,
•				35695917, 35695855, 264635, 264636,
				83373044, 264486
38094948 (5195,	88094948 (5195, 5196) Novel Protein sim. GBank gi[1001351 dbj BAA10838  -		UNCLASSIFIED	22278998, 264259, 29331824, 87168474,
•	(D64006) hypothetical protein (Synechocystis sp.)			264683, 21906766, 35695917, 264691,
				33657023, 33657109, 18108370, 18108374,
				264564, 264565
37642889 (5187.	87642889 (5187, 5198) Novel Protein sim. GBank gij3941737 (AF109719) - BAT2		MHC	264766, 264769, 21906768, 33657182,
	[Mus musculus]			35695763, 18108370, 18108374, 264635,
				204030, 30320400, 22273000, 204300
37787846 (5199.	87787846 (5199, 5200) Novel Protein sim. GBank gil4263521lgb AAD153471-	Contains protein domain (PF00400) - kinasereceptor	cinasereceptor	35696286, 264093, 264288, 21906769,
	(AC004044) putative WD-repeat protein [Arabidopsis	WD domain, G-beta repeat		35696423, 35695855
	Inaliana			

100	04040030 45004 500	1 20 11 11 11	10.000	EC101575 22320000 264260 20221922
	91243070 (3201, 321	Zoo!   Pizesuru (azor, azoz) novel Protein sim. Gbank	DODGE STATE OF THE	29331824, 60432289, 29331826, 29331827.
		SQ WARNING ENTRY III		264906, 265007, 265008, 264591, 60433356,
				33657402, 60433438, 21906754, 265011,
				265018, 265019, 18108351, 264448, 264369,
				21906769, 265020, 60170615, 264693,
				33657109, 18108370, 18108376, 56182323,
				18108381, 18108385, 22279002, 264563
2602	88180022 (5203, 5204)	Novel Protein sim. GBank gi 4406632 gb AAD20047  -		60433438, 21906754, 87168559, 264601.
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			•	265009, 60433356, 264757, 60433438,
				264759, 33657084, 87168474, 265010,
				265011, 87168559, 265017, 265018, 265019,
				264448, 264683, 18108354, 264288, 264767,
				264689, 21906765, 21906766, 21906767,
			سيسي	21906768, 21906769, 55811957, 265020,
			-	265021, 265022, 60170615, 264691,
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	-			18108374, 35696423, 65274791, 35695855.
				264555, 264636, 264637, 56182323,
				83373044, 56526486, 87168518, 60432113,
				22279000
2604	94676601 (5207, 520	94676601 (5207, 5208) Novel Protein sim. GBank	ancogene	264259, 35696052, 264508, 264906, 264907,
		gij5454030jretjNP_006468.1jpRRP2 - RAS-related on		264908, 264909, 264510, 264512, 265008,
				264910, 33657402, 264604, 264605, 264762,
				264763, 264682, 264764, 264683, 264768,
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				264639, 83373044, 264565
2605			UNCLASSIFIED	22278998, 264490, 60432049, 264259,
	-	(AB013721) mitsugumin 23 [Oryctolagus cuniculus]		60432289, 264909, 265008, 60433356,
			_	60433438, 264758, 21906754, 265010,
				265011, 265018, 264681, 18108351, 264288,
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9000	07746400 /6044 604			22272006 264510 284512 285000 264758
9097	87745405 (5211, 5212)	(7)		22279002, 264566
			1	2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2007, 2

29331822, 29331825, 29331826, 29331827, 29331828, 264906, 264907, 264908, 66712502, 264828, 56182435, 55812038, 265010, 265017, 265018, 265019, 264768, 265020, 26692, 33657023, 264693, 33657023, 264693, 33657109, 18108370, 264639, 56182323, 264630, 264590, 264590, 264590, 264590, 264590, 264592, 264758, 264590, 264590, 264592, 264758, 264590, 264590, 264592, 264758, 264590, 264590, 264592, 264758, 264590, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264590, 264592, 264758, 264590, 264592, 264758, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264500, 264590, 264590, 264590, 264590, 264590, 264590, 264590, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 264500, 26450	264631 254488, 65274572, 35696286, 22278997, 22278999, 60432049, 264259, 56182181, 29331824, 29331825, 69331828, 60432289, 29331827, 29331828, 264905, 264907, 60433356, 60433438, 55812038, 265011, 87168559, 265017, 265018, 264448, 264765, 264288, 264766, 264699, 21906765, 21906767, 21906769, 285020, 265021, 264691, 264692, 33657109, 27486261, 18108370, 65274791, 264636, 264556, 56182323, 18108385, 56526486	18108394, 22278994, 56994075, 60432049, 264259, 29331822, 29331825, 60432289, 29331827, 264109, 264905, 56182435, 264102, 265006, 265007, 265009, 60433356, 60433438, 265011, 87168559, 265017, 264448, 264682, 264764, 264288, 265021, 33657023, 263967, 33657182, 27486261, 18108374, 263976, 55811576, 264638, 87168518, 60432113	22278999, 265017, 264684, 21906768, 22279000	265009, 264910, 264759, 265017, 21906/67, 18108365, 18108388, 60432113	264685	22278998, 22278999, 29331825, 264508, 264906, 21906754, 264602, 264766, 264769, 52644229, 21906765, 33657109, 27486264, 18108370, 263972, 264555, 60432113	264768, 18108394, 264592, 264693, 264508, 264509, 264509, 264907, 264628, 264908, 264909, 18108377, 264511, 264512, 264910, 264595, 265010, 264404, 264563, 264764, 264685, 264766
	histone	transcriptfactor	isomerase	transport	UNCLASSIFIED		UNCLASSIFIED
	Contains protein domain (PF00850) - histone Histone deacetylase family		Contains protein domain (PF00254) - isomerase FKBP-type peptidyl-prolyl cis-trans Isomerases		and the second s	Contains protein domain (PF00625) - Guanylate kinase	
2607       87627742 (5213, 5214)   Novel Protein sim. GBank gil4826626 gb AAD30202.1  -         (AF135022) mediator [Homo saplens]         (AF135022) mediator [Homo saplens]         2608       81734786 (5215, 5216)   Novel Protein sim. GBank gil22266005 (U49973) - ORF2:		88177654 (5219, 5220) Novel Protein sim. GBank gil4336855lgblAAD17989] - (AF106473) teucine-rich-domain inter-acting protein 1; LEAP1 [Mus musculus] inter-acting protein 1; LEAP1 [Mus musculus]	87428890 (5221, 5222) Novel Protein sim. GBank gi[3876761 emb CAA92994  - Contains pre (Z68760) predicted using Genefinder; Similarity to Mouse FKBP-type I FK506-binding protein (SW:FKB3_MOUSE) [Caenorhabditis isomerases elegans]	87771198 (5223, 5224) Novel Protein sim. GBank gil5679136]gb AAD46874.1 AF16093 - (AF160934) BCDNA_LD14189 [Drosophila melanogaster]		87643948 (5227, 5228) Novel Protein sim. GBank gi 5533081 gb AAD45009.1 AF16118 - (AF161181) P55T protein [Mus musculus]	
87627742 (5213, 5214) 81734786 (5215, 5216)	94843791 (5217, 5218)	88177654 (5219, 5220)	87428890 (5221, 5222)	87771198 (5223, 5224)		87643948 (5227, 5228)	87381996 (5229, 5230)
2607	2609	2610	2611	2612	2613	2614	2615

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	predicted using Ge			22278999, 264259, 29331822, 29331824,
	elegans			56182181, 29331825, 60432289, 29331828,   35696052, 29146499, 66712502, 52644045
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8 (5249, 5250)	86452068 (5249, 5250) Novel Protein sim. GBank gij2887429 dbjjBAA24857  -	=	UNCLASSIFIED	264091, 264511, 263981
	(AB007887) KIAA0427 [Homo sapiens]			
37 (5251, 5252)	16533797 (5251, 5252) Novel Protein sim. GBank gi]487416 (L20302) - actin		struct	265008
	filament protein [Gallus gallus]			
23 (5253, 5254)	87636823 (5253, 5254) Novel Protein sim. GBank gij88462 pir JA27307 - proline-rich	=-	UNC! ASSIFIED	22278996, 265007, 265009, 264448,
	phosphoprotein (gene PRH1, Db allele) - human			21906767, 265021, 264558, 87168518
54 (5255, 5256)	94848254 (5255, 5256) Novel Protein sim. GBank gij3123552jembjCAA18609j -		UNCLASSIFIED	22278997, 22278999, 264259, 60432049,
	(AL022578) dJ393P12.2 (hypothetical Proline-rich protein			29331822, 29331824, 29331825, 29331827,
	KIAA0269 LIKE) [Homo sapiens]			35696052, 29331828, 264907, 264909,
				265008, 264591, 60433356, 60433438,
				265010, 265017, 265018, 264369, 264288,
				18108357, 21906765, 21906768, 265022,
				65274791, 264638, 18108387, 87168518,
				22279002
90 (5257, 5258)			synthase	29331825, 29331826, 264102, 265006,
	gi 4929595 gb AAD34058.1 AF15182 - (AF151821) CGI-63			264766, 35695917, 264691, 33657023,
	protein (Homo sapiens)			263972, 18108374, 22279000
79188364 (5259, 5260)		)	UNCLASSIFIED	264636, 18108385
09 (5261, 5262)	305[pir]JQ1161 - Gag	Contains protein domain (PF00098) - c	Ina_rna_bind	52644507, 52645156, 52646365, 52646842,
	(V1)	Zinc finger, CCHC class		22278994, 22278995, 35696286, 56994075,
				22278997, 22278999, 264259, 52645080,
				29331822, 29331824, 29331825, 29331826,
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				33657182, 27486261, 27486262, 27486265.
				33657349, 35695763, 35696423, 65274791,
				35695855, 264634, 264637, 52644332,
				56182323, 60432113, 264566, 264486
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22276995, 22278997, 22278998, 264259, 29331822, 29331824, 29331824, 29331825, 29331827, 264508, 264908, 265006, 265007, 265008, 265009, 265010, 265011, 87168559, 265018, 265019, 264683, 264288, 264685, 29148629, 33657023, 264693, 33657182, 35693763, 55811578, 264693, 33657182, 35693763, 18108385, 56526486, 87168518, 22279000, 22279002, 264565	29331826, 263972, 264089	22278997, 264259, 29331824, 66714117, 29331827, 29331828, 264907, 33657084, 265017, 265018, 264448, 264288, 21906766, 21906767, 21906768, 29148629, 18108376, 55811576, 35695855, 87168518, 22279000	264569, 29331822, 29331828, 265006. 60170831, 264681, 264765, 264685, 29148627, 21906769, 29148784, 265022, 60170615, 264635, 18108385, 56526486, 22279002, 264567	264488, 264489, 52644507, 264887, 52646365, 52646842, 22278994, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 20281171, 264259, 29331822, 29331827, 35696052, 29331828, 29146498, 264508, 264905, 264905, 264907, 264907, 264909, 264511, 264512, 264910, 264591, 264592, 264593, 264593, 264603, 264605, 264761, 264600, 264602, 264603, 264764, 264765, 264769, 21906765, 21906765, 21906765, 21906766, 21906767, 21906765, 21906769, 21906765, 21906763, 264691, 356524, 18108374, 35696423, 264628, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264657, 264667, 264466, 264466, 264567
	UNCLASSIFIED			phosphatase
Contains protein domain (PF00084) - Sushi domain (SCR repeat)			Contains protein domain (PF01546) - Peptidase family M20/M25/M40	Contains protein domain (PF00782) - phosphatase Dual specificity phosphatase, catalytic domain
2633   95011617 (5265, 5266) Novel Protein stm. GBank gil1139548 db  BAA10889  - (D84009) seizure-related gene product 6 type 2 precursor [Mus musculus]	87330921 (5267, 5268) Novel Protein sim. GBank gil54416111emb CAB46854.1 - (AJ388555) hypothetical protein [Canis familiaris]	86623144 (5269, 5270) Novel Protein sim. GBank gil4680663]gb AAD27721.1 AF13294 - (AF132946) CGI-12 protein [Homo sapiens]	87260534 (5271, 5272) Novel Protein sim. GBank gij3879146 emb CAB07646  - (293386) Similarity to Yeast hypothetical 52.9 KD protein (SW:P43616); cDNA EST EMBL:M89432 comes from this gene; cDNA EST EMBL:D71008 comes from this gene; cDNA EST EMBL:D73578 comes from this gene; cDNA EST EMBL:D73578 comes from this gene; cDNA EST EMBL:D69025 comes from this gene; cDNA	95011299 (5273, 5274) Novel Protein sim. GBank gil4759208[ref NP_004081.1 pDUSP - dual specificity phosphatase 3 (vaccinia virus phosphatase VH1-related)
95011617 (5265, 5266)	$\overline{}$	86623144 (5269, 5270)	87260534 (5271, 5272)	95011299 (5273, 5274)
263.	2634	2635	2636	2637

<u> </u>	gil4829689[gb[AAD34105.1 AF15186 - (AF151868] CGI-110 protein [Homo sapiens]	AF15186 - (AF151868) CGI-110 KNA recognition motif. (a. K.a. Kraw, RBD, or RNP domain)		50 (1025) 7, 242 (10393), 245 (10394), 250 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030) (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030 (2030) (2030) (2030 (2030) (2030 (2030) (2030 (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2030) (2
7, 5278) NK (A 9ri	95361346 (5277, 5278) Novel Protein sim. GBank gi[2190007 dbj BAA20355  - (AB004109) phosphatidylserine synthase II [Cricetulus griseus]	, vi	synthase	264488, 29331825, 35696052, 264508, 264509, 264909, 264512, 33657402, 60433438, 264758, 85658542, 264600, 265020, 265021, 33657109, 264628, 35696423, 264555, 264639, 264563, 264564, 264565, 264566, 264486
9, 5280) Nr. CG 27 (P) (P)	87781330 (5279, 5280) Novel Protein sim. GBank gil3158516 (AF067617) - contains similarity to chromo (chromatin organization modifier) domains (Pfam: chromo.hmn., score: 17.76 and 27.94) and to helicases conserved C-terminal domain (Pfam: helicase_C.hmm, score: 67.00) [Caenorhabditis etegans]	Contains protein domain (PF00271) - helicase Helicases conserved C-terminal domain	elicase	29331822, 29331826, 264906, 33109954, 265017, 265019, 21906768, 35695763, 264636, 264637, 18108387
11669834 (5281, 5282)			UNCLASSIFIED	264828
3, 5284) Ni	87412575 (5283, 5284) Novel Protein sim. GBank gil2564955 (AF030001) - unknown [Mus musculus]		UNCLASSIFIED	264259, 29331822, 33657402, 265019, 264369, 264691, 264634, 56526486, 22279002
5, 5286) N. (A	87643961 (5285, 5286) Novel Protein sim. GBank gil4490304 emb[CAB38795.1] - (AL035678) putative protein [Arabidopsis thaliana]	Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase	ıelicase	22278997, 264259, 29146499, 56182435, 264910, 265010, 18108351, 264682, 284663, 264369, 264686, 29148627, 264690, 33657109, 18108370, 263973, 18108374, 264684, 264557, 264558,
7, 5288) Ni (A	88177671 (5287, 5288) Novel Protein sim. GBank gil3769797[gb]AAC67502.1] - (AF059569) actin binding protein MAYVEN [Homo sapiens]	Contains protein domain (PF00651) - Inucl_recpt BTB/POZ domain	nucl_recpt	264107, 264687
9, 5290)	17277228 (5289, 5290)	1		265007
1, 5292) N 9i St	Novel Protein sim. GBank gi 1706722 sp P49749 EVX2_MOUSE - HOMEOBOX EVEN SKIPPED HOMOLOG PROTEIN 2 (EVX-2)		UNCLASSIFIED	264909, 264687, 264632, 83373044

2647	2647 91212978 (5293, 5294)			CULLIA CONTIN		ſ
				discossing the second s	Ja 1023/2, 22278998, J2695286, 22278998, 264259, 29331822, 56182181, 29331825,	
					60424269, 60432289, 35696052, 66712502,	
					264908, 265007, 55812038, 33109954, 21006254, 2265264, 26564	
					21900/34,3383/084,263019,264448, 264388 66484663 34886466 34886436	
					274255, 30181352, 21808753, 21808756, 21908768, 21908769, 35695917, 365030	-
				_	265021 42644150 264603 22652400	
				•	200021, 32044130, 204033, 3383/109, 33657340, 60424629, 4640634, 7564020	
					35001349, 00431328, 18108374, 35810764,	
		-		4	33090423, 36182323, 60432113, 22279002, 264664	
2648	87600587 (5295, 5296)				204304 20146408 56182435 33108064 266044	Т
				ir -	23140430, 30102433, 33103834, 263011, 264682 55811957 35695017 364600	_
95	244004 1007, 1000				263976, 18108377, 35696423, 60432113	
25	94120,03 (3287, 3288)	34 120/03 (323/, 3288) Novel Protein Sim. GBank gi 3041852 (AC004539) -		UNCLASSIFIED	56182575, 35696286, 22278998, 29331824	<u> </u>
		unknown runction; similar to Y09105 (PID:g1666171) [Homo  sanians			29331826, 60432289, 66712502, 56182435,	_
					60170831, 60432229, 33657402, 33109954,	_
					21906754, 265017, 264686, 264688,	
					21906765, 21906768, 60170615, 264693,	
					263967, 18108370, 263976, 60170394,	
2650	87297533 (5299 5300)	87297533 (5209 5300) Novial Dratain aim CBant - 11525533 1111			60432113, 22279002, 264563	
	(000, 000)	Novel Floteill Sim. Chank gijosboz77   dbj BAA81908.1   -   (AB029335) HrPET-3 (Halocynthia rozetzi)			264685	_
2651	88088745 (5301, 5302)	88088745 (5301, 5302) Novel Protein sim. GBank gil4240225IdhilRAA74891 11.	Contains activities demain (DESCOPES)			_
		(AB020675) KIAA0868 protein (Homo sapiens)	Contains protein domain (Produs4) - synthase Laminin G domain		29331824, 29331826, 29331827, 265007, 55843028, 34006754, 4846600, 4846600	
_	30000				33812038, 21908/34, 18108366, 18108384, 22279002, 264567	
7007	10343123 (3303, 3304)			UNCLASSIFIED	264692	Υ-
	87798735 (5305, 5306)	8/798/35 (5305, 5306) Novel Protein sim. GBank gil4493956jemb CAB11123.2] -		Γ	265018 18108370 18108387 264566	_
		(Z98551) predicted using hexExon; MAL3P6.28	-		200000000000000000000000000000000000000	
		(PFC0845c), Hypothetical protein, Ien: 167 aa; Similarity to				_
		model organism hypothetical proteins (C.elegans,				
		D.melanogaster, S.cerevisiae & S.pombe). C.elegans protein ZK287.5 (TR				
2654	95103240 (5307, 5308)				60424179 65274572 56182575 364250	<del></del>
					56182181 264908 56182434 55814067	
					35695917, 265021, 263976, 55810764,	
2655	91229018 (5309 5310)	91279018 (5309 5310) Mariel Protein Sim CB. 1. 1977577			65274791, 56182323, 83373044, 65274727	
	(1010) (2000) (2010)	Novel Florein sim. Gbank gij36/52/2 embjCAB02861  -	Contains protein domain (PF00097) - transcriptfactor		56182575, 56181686, 264092, 264259,	τ
_		C3HC4 type (RING finer): CONA EST WAARDS 3 comes	Zinc finger, C3HC4 type (RING		56182181, 60432289, 264907, 33657402,	
		from this gene; cDNA EST vk443h5 5 comes from this cene	(labin)		55812038, 21906754, 87168559, 265017.	_
		[Caenorhabditis elegans]			264448, 264369, 264288, 21906765,	_
	<u> </u>				21906/66, 21906/67, 21906/68, 3365/109, 18108370, 264628, 65844676, 264666	
					264639, 83373044, 56526486, 264404,	
2656	84562601 (5311, 5312) N	84562601 (5311, 5312) Novel Protein sim GBank oil3043718IdhilBa & 255231			60432113	
	)				264693	

2657	52561728 (5313, 5314)	52561728 (5313, 5314) Novel Protein sim. GBank gil5689509 dbj BAA83038.1  -   ABA79609 KIAA1086 protein Homo sanjasel		dna_rna_bind	264693
2658	88062454 (5315, 5316)	88062454 (5315, 5316) Novel Protein sim. GBank gij3686089 (AC005757) - R32611_1 [Homo sapiens]	Contains protein domain (PF00560) - nucleaseinhib Leucine Rich Repeat	nucleaseinhib	35696286, 264259, 29331822, 29331824, 29331826, 29331828, 295019, 264683, 21906768, 35695917, 264693, 35695855, 264637, 87168518, 264486, 264567
2659	87600755 (5317, 5318)	87600755 (5317, 5318) Novel Protein sim. GBank gij5420387jembJCAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]	Contains protein domain (PF01426) - UNCLASSIFIED BAH domain	UNCLASSIFIED	264909, 264910, 265018, 264369, 264769, 21906769, 264693, 263972, 18108388
2660	-	91718472 (5319, 5320) Novel Protein sim. GBank	Contains protein domain (PF00036) - kinase	kinase	264488, 65274572, 35696286, 22278998,
		gij728837 sp P39194 ALU7_HUMAN - !!!! ALU SUBFAMILY   EF hand	EF hand		22278999, 264259, 29331822, 29331824,
		SQ WARNING ENTRY !!!!			60432289, 29331826, 35696052, 264908.
					56182435, 265008, 265009, 60433356,
					264594, 265010, 265018, 55811150,
					18108351, 264682, 264684, 264369, 264288.
					264687, 21906765, 29148784, 35695917,
					001/0015, 52644150, 5365/025, 5565/103, 35696423, 35695855, 264556, 60170394
					18108385, 22279000, 22279002
2661	95342817 (5321, 5322)	95342817 (5321, 5322) Novel Protein sim. GBank		glycoprotein	60432049, 264259, 29331824, 29331825,
		gil4758048/refINP 004739.1 pCPR8 - cell cycle progression			29331826, 29331827, 29331828, 264906,
		8 protein	,		264909, 264593, 33109954, 265010, 265017.
					265018, 265019, 264760, 264448, 264369,
					264288, 21906765, 21906768, 265022,
					264691, 33657023, 27486262, 60431528.
					18108374, 35695855, 18108388, 264482
2992					264555, 264556, 264558, 264486
2663	_	87780623 (5325, 5326) Novel Protein sim. GBank gij3874714 emb[CAA91263] -		dehydrogenase	264906, 264909, 264757, 264758, 264767,
		(Z66494) similar to choline dehydrogenase; cDNA EST			264691, 33657023, 264638
		yk346d5.5 comes from this gene; cDNA EST yk346d5.3			
7990	06640000 (6007 6000)	comes from this gene (Caenornabditis elegans)	(900000) cicmob cichose pointer	0000000	35606386 364500 364505 364388 364685
007	02210258 (2327, 2320)	Nover   Flucial State   Code   Policial   Notice   Notice     Notice   No	Contains protein contain (Frocos) - EGF-like domain	auañonio	264686
2665	•	87770662 (5329, 5330) Novel Protein sim. GBank gij4884406jemb CAB43311.1 -		UNCLASSIFIED	35696286, 22278999, 29331822, 35696052,
		(AL050190) hypothetical protein [Homo sapiens]			264906, 264907, 264909, 264510, 264511,
					264512, 264593, 60433438, 265019, 264681.
					21906765, 21906766, 21906767, 21906768,
					265020, 265022, 35696423, 35695855,
	_				22279002, 264482, 264486
2666		87826472 (5331, 5332) Novel Protein sim. GBank		UNCLASSIFIED	29331825, 265007, 264910, 60432229.
		gi[5106956]gb[AAD39906.1]AF11361 - (AF113615) EH1/EH2 domain-containing stollain EUOS (Homo service)			265019, 264288, 21906767, 264558, 2227ann2
2667		R7422720 (5222 5224) March Decision aim Charle	Contains associated and the Contain (DEC)	eacolone	264007 20111810 264681 264681 264288
2007		NOVEL PLOTEIN SITH, GBAILK	2. contains protein domain (r.r.d. 130) -	aceanni	104301, 2331030, 204001, 204003, 204200, 104200, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 1043000, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 1043000, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 104300, 1043000, 1043000, 1043000, 1043000, 1043000, 1043000, 1043000, 10430000, 1043000, 1043000, 1043000, 1043000, 1043000, 10430000, 10430000, 1
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UNCLASSIFIED			transport	- UNCLASSIFIED
		Contains protein domain (PF00628) PHD-finger		Contains protein domain (PF01344) - UNCLASSIFIED Ketch motif
2668   91216716 (5335, 5336) Novel Protein sim. GBank   gil\$4\$4186 ref NP_006327.1 pZYG  - ZYG homolog	95415721 (5337, 5338)       Novel Protein sim. GBank gi[2147012 pir] JC4899 - proline         rich protein - rat       rat	87613234 (5339, 5340) Novel Protein sim. GBank gij1723523 sp Q10362 yDBB_SCHPO - HYPOTHETICAL 94.9 KD PROTEIN C22E12.11C IN CHROMOSOME I	91214936 (5341, 5342) Novel Protein sim. GBank gil4768277[gblpAAD29444.1JAF06425 - (AF064255) very long-chain acyl-CoA synthetase homolog 2: VLCS-H2 [Homo sapiens]	2672 87399123 (5343, 5344) Novel Protein sim. GBank gil4966346 gplAAD34677.1 AC00634 - (AC006341) Contains two PF 01344 Kelch motif domains.  Arabidopsis thaliana]
91216716 (5335, 5336)	2669   95415721 (5337, 5338)	2670 87613234 (5339, 5340)	<u>2671   91214936 (5341, 5342)</u>	72 87399123 (5343, 5344

otein sim. GBank  24 ref NP_004280.1 pNRF3 - nuclear factor  4-derived 2)-like 3  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  AMP-binding enzyme	30/49 (3545, 3545)	2873   87430749 (5345, 5346) Novel Protein sim. GBank gil5457337 emb CAB41505.2  - (AJ236876) poly(ADP-ribosyl) polymerase-2 (Homo sapiens]	Contains protein domain (PF00644) - polymerase Poly(ADP-ribose) polymerase catalytic region	polymerase	22278995, 22278996, 22278997, 2227899, 29331822, 29331824, 29331828, 35696052, 60434498
Contains protein domain (PF00170) - transcriptfactor  DZIP transcription factor  DZIP transcription factor  DZIP transcription factor  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  Sophila melanogaster)  AMP-binding enzyme					265018, 265019, 26448, 21906768, 2185018, 265019, 26448, 21906768, 21906769, 265020, 265021, 33657109, 27466767, 24666767, 6443467, 644370304
Contains protein domain (PF00170) - transcriptfactor bZIP transcription factor  UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED OSophila melanogaster)  AMP-binding enzyme					27460518, 264563
IpNRF3 - nuclear factor bZIP transcription factor  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  Sophila melanogaster) AMP-binding enzyme	(5347, 5348,		Contains protein domain (PF00170) -	transcriptfactor	264488, 22278996, 35696286, 264091,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED OSophila melanogaster) AMP-binding enzyme		₹	bZIP transcription factor		264259, 29331824, 29331826, 35696052,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED OSophila melanogaster) AMP-binding enzyme		(erythroid-derived 2)-like 3			264511, 55812038, 85658542, 264766,
UNCLASSIFIED UNCLASSIFIED UNCLASSIFIED Osophila melanogaster) AMP-binding enzyme AMP-binding enzyme					Z1906/65, 35695917, Z646Z9, 356964Z3, 18108383, 87168518
UNCLASSIFIED 1079042 pir  S52154 - acetyl- Contains protein domain (PF00501) - synthase 0sophila melanogaster) AMP-binding enzyme	(5349, 5350)			UNCLASSIFIED	264691
1079042 pir  S52154 - acetyl- Contains protein domain (PF00501) - synthase osophila melanogaster) AMP-binding enzyme	(5351, 5352)			UNCLASSIFIED	264906, 265008
AMP-binding enzyme	(5353, 5354)	1079042 pir  S52154 - acetyl-	Contains protein domain (PF00501) -	synthase	18108394, 65274572, 56182575, 22278994,
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					265008, 264910, 265009, 264593, 264594,	
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					21906754, 265010, 265011, 87168559,	
					264600, 264601, 264602, 264603, 265017.	
					265019, 264605, 264760, 264762, 18108351,	
					264763, 264682, 264683, 264764, 264288,	
					264766, 264687, 264768, 264769, 264689,	_
					21906765, 21906767, 35695917, 265020.	
					265021, 52644150, 264691, 33657023,	_
_					264693, 18108364, 18108365, 33657109.	_
_					33657349, 264628, 18108374, 35696423,	
		-			35695855, 264630, 264631, 264632, 264634,	_
					264635, 264555, 264636, 264637, 264638,	_
					264639, 18108385, 264563, 264564, 264565,	
					264566, 264486, 264567	
2679	95361544 (5357, 5358)	95361544 (5357, 5358) Novel Protein sim. GBank	Contains protein domain (PF00970) - reductase	reductase	264488, 22278996, 35696286, 264259,	Τ
		git709233IspiP07514INC5R BOVIN - NADH-	FAD/NAD-binding Cytochrome		29331826, 29331827, 29331828, 264909,	_
		CYTOCHROME BS REDUCTASE	reductase		56182435, 264113, 264511, 265008,	
					60433438, 264758, 85658542, 87168474.	_
					265011, 265017, 265019, 264288, 21906766,	-
					21906767 21906768 21906769 55811957	_
					265021, 264690, 33657023, 55810764.	_
					35696423 55811576 264631 18108381	_
					550504£3, 55011576, £04551, 10150551, 160170394, 83373044, 87168518, 264566	_
2680	87800356 (5359 5360)	RTROUTSE (5350 5350) Navel Protein sim CBank nid 58060Aldhil BAA76804 11	Contains protein domain (DE0003B)	etrict	264259 264102 264905 264908 265007	Т
	de cocco (cocco), cocco	NOVELT FOREIT SHIP: OBSIDE 9143589004[db] BXX1,0024.11 -	FF hand		204209, 204102, 204900, 204300, 205001, 1265008, 33109954, 265011, 18108351	
					264768 33657023 20281149 263972	_
					264630, 264635, 264638	
2681	90933844 (5361, 5362)	90933844 (5361, 5362) Novel Protein sim. GBank		kinase	264489, 22278995, 264509, 264905, 264906,	
	•	gil728837 splP39194 ALU7 HUMAN - !!!! ALU SUBFAMILY			264907, 264908, 264909, 264510, 265006,	
		SQ WARNING ENTRY IIII			264512, 264910, 264594, 60433438, 264758,	
					264603, 264604, 265019, 264605, 264760,	_
					264764, 264687, 264768, 264769, 21906769,	
_					35695917, 265020, 33657023, 264631,	
					264635, 264637, 264638, 264639, 264486	
2682	94138934 (5363, 5364)	94138934 (5363, 5364) Novel Protein sim. GBank gil423468 pir JQ1974 - HTF9-C	Contains protein domain (PF00076) - (	UNCLASSIFIED	35696286, 264908, 55811386, 265017,	
			RNA recognition motif. (a.k.a. RRM,		55811150, 55811957, 35695917, 60431528,	
			RBD, or RNP domain)		55810764, 55811576, 35696423, 65274791,	
					56526486	Т
2883	87774405 (5365, 5366)	87774405 (5365, 5366) Novel Protein sim. GBank gi[5114351[gb]AAD40286.1[-	Contains protein domain (PF00622) - interleukinrecept	interleukinrecept	264909, 264769, 264635, 264636	
		ואוא נו ישני ון אווים ווושנים וושנים ושלווו סאוא נו ישני און	SPRT domain			٦

-1					
2684	85787151 (5367, 5368)	85787151 (5367, 5368) Novel Protein sim. GBank gil4886469 emb CAB43385.1  [(AL050284) hypothetical protein [Homo sapiens]			264593
2685	88054299 (5369, 5370)	88054299 (5369, 5370) Novel Protein sim. GBank gij3342729 (AC005331) - R31341_2 [Homo sapiens]		UNCLASSIFIED	
2686		87628690 (5371, 5372) Novel Protein sim. GBank gil4650844 db  BAA77027.1  - (AB026190) Kelch motif containing protein [Homo sapiens]	Contains protein domain (PF01344) - struct Kelch motif		264259, 29331822, 60432289, 29331827, 29331830, 264909, 264512, 264596, 264769, 264534, 264555, 264556, 264557, 264558, 60170394, 264559, 264486
2687	87998183 (5373, 5374)	87898183 (5373, 5374) Novel Protein sim. GBank gi 5281314 gb AAD41475.1 AF13312 - (AF133123) transcription factor IIIC102 [Homo sapiens]	Contains protein domain (PF00515) - transcriptfactor TPR Domain		18108394, 18108396, 22278996, 35696286, 22278997, 29331826, 29331828, 66712502, 21906754, 265011, 264760, 264761, 264763, 264689, 21906765, 35696423, 264559, 18108385, 264563
2688	79959584 (5375, 5376)				264908, 264760
2689	94122440 (5377, 5378)	94122440 (5377, 5378) Novel Protein sim. GBank gi 3880023 emb CAA97339  - (Z73098) Similarity to yeast hypothetical protein (Swiss Protacession number Q09695); cDNA EST EMBL:D72982 comes from this gene; cDNA EST EMBL:D75947 comes from this gene [Caenorhabditis elegans]			22278997, 22278998, 22278999, 29331824, 35696052, 264908, 264908, 56182435, 264910, 265009, 60433438, 21906754, 18108351, 264682, 264863, 264767, 21906765, 21906765, 21906768, 33657023, 33657182, 27486262, 27486264, 33657349, 18108370, 18108372, 18108374, 35696423, 3569555, 18108385, 22279002
2690		88003055 (5379, 5380) Novel Protein sim. GBank gi[2477513 (AC002398) - [F25965_3 [Homo sapiens]	Contains protein domain (PF00620) - struct RhoGAP domain	struct	
2691	91219241 (5381, 5382)	91219241 (5381, 5382) Novel Protein sim. GBank gil4107276 emb CAA67130  - (X98506) acetyl-CoA synthetase [Solanum tuberosum]		synthase	65274572, 56182575, 264259, 29331822, 29331824, 29331826, 29331828, 264112, 265009, 55812038, 264596, 33109954, 265017, 264448, 264288, 264369, 264684, 21906769, 60170615, 60431528, 55810764, 264634, 264636, 264556, 264637, 22279002, 264566, 264566
2692	94111914 (5383, 5384)	94111914 (5383, 5384) Novel Protein sim. GBank gij3513303 (AC005594) - R26984 1 (Homo sapiens)	Contains protein domain (PF00326) - peptidase Protyl oligopeptidase family	peptidase	
2693	20438807 (5385, 5386)			UNCLASSIFIED	264592
2694	T	94111918 (5387, 5388) Novel Protein sim. GBank gij3122400[spl035682]MUG_MOUSE - MYELOID UPREGULATED PROTEIN			264559
2695		Novel Protein sim. GBank gil4972740[gb]AAD34765.1  - (AF132177) unknown [Drosophila melanogaster]		collagen	35696286, 56994075, 22278999, 264259, 35696052, 29331830, 265011, 264288, 56181562, 264690, 264692, 33657023, 27486262, 263976, 18108376, 35696423, 35695855, 60170394, 83373044, 55525486, 22279000, 22279000, 264566
2696	87874040 (5391, 5392)	87874040 (5391, 5392) Novel Protein sim. GBank gij728831 sp P39188 ALU1_HUMAN - III! ALU SUBFAMILY J WARNING ENTRY III!		synthase	264594, 21906768, 18108370, 18108372

35696286, 264259, 29331824, 29331825, 29331826, 29331826, 29331827, 29331828, 35696052, 265006, 265007, 264512, 265009, 60170831, 60433356, 264595, 284788, 81768474, 265010, 265011, 8176859, 264601, 265017, 264288, 264766, 264688, 264689, 21906765, 21906768, 265020, 265021, 60170615, 33657109, 18108376, 35696423, 3569565, 264555, 264556, 6017034, 244559, 18108387, 56526486, 87168518, 22279002, 264553, 284656	22278995, 35696286, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 29331825, 60432289, 29331822, 29331827, 33656970, 264906, 29331830, 264909, 52644045, 264910, 60433356, 33657402, 33109954, 265017, 265018, 265019, 264288, 21906765, 21906766, 21906767, 21906769, 231696370, 265021, 265021, 265022, 52644150, 33657023, 33657182, 27486261, 35696423, 65274791, 284638, 60432113, 22279000	264768, 18108357, 264690, 264691	60424269, 56182435, 60432229, 60433438, 55811386, 265017, 55811150, 264448. 56181562, 55811957, 264693, 33657109, 60431528, 264629, 55810764, 55811576, 65274791, 60431850, 60432113	18108394, 52645156, 35696286, 264259, 29331822, 29331824, 29331826, 60432289, 29331826, 29331827, 35696052, 29331828, 29146499, 265006, 60433356, 33657402, 60433438, 264595, 33657084, 18108351, 264288, 264769, 18108359, 21906768, 35659517, 33657023, 27486261, 18108374, 18108388, 60432113, 22279000	29331828, 264512, 264555, 264556, 264557, 264558, 264559	60432289, 265007, 21906765, 21906768, 265021, 264563
UNCLASSIFIED	Iranថ្មីឆ្នាំpifactor	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	
	Contains protein domain (PF00412) - tran্রীক্রাptfactor LIM domain containing proteins			-		Contains protein domain (PF00642) - Zinc finger C-x8-C-x5-C-x3-H type (and similar).
91638472 (5393, 5394) Novel Protein sim. GBank gi[5689473 dbj BAA83020.1  - (AB028991) KIAA1068 protein [Homo sapiens]	94325891 (5395, 5396) Novel Protein sim. GBank gil841318 (U22818) - mulant sterol regulatory element binding protein-2 (Cricetulus griseus)	87780650 (5397, 5398)	Novel Protein sim. GBank gi 5174395 ref NP_006006.1 pB120 - Brain protein 120	94148584 (5401, 5402) Novel Protein sim. GBank gil1572801 (U70854) - F38A5.1 gene product [Caenorhabditis elegans]	57295366 (5403, 5404) Novel Protein sim. GBank gi[2605967 (AF030027) - 24 [Equine herpesvirus 4]	87649514 (5405, 5406) Novel Protein sim. GBank gi 5689399 dbj BAA82983.1  - (AB028954) KIAA1031 protein [Homo sapiens]
91638472 (5393, 5394)		87780650 (5397, 5398)	94139836 (5399, 5400)			
2697	2698	2699	2700	2701	2702	2703

2704	87649515 (5407, 5408)	2704 [87849515 (5407, 5408)]Novel Protein sim. GBank oil4335694inhIAAB632941.		264488 22278995 22278998 20231828
		(AF008554) Implantation-associated protein [Rattus		29146499, 264905, 264906, 264907,
		norvegicus]		52644045, 264511, 33657402, 264600,
				264602, 265017, 264605, 264761, 18108351,
				264764, 264687, 264769, 265021, 264691,
				264692, 18108362, 264693, 18108370,
				18108374, 264634, 264835
2705	87771745 (5409, 5410)			264489, 264509, 264511, 264512, 264910,
				264593, 87168474, 264604, 264288, 264687,
				264769, 264638, 264566, 264486
2706		94326789 (5411, 5412) Novel Protein sim. GBank gij3255952 emb CAA16821.1	UNCLASSIFIED	264488, 52646842, 65274572, 22278994.
		(AL021728) /prediction=(method:; /match=(desc:		56994075, 22278997, 264259, 29331824.
		[Drosophila melanogaster]		29331825, 29331826, 29331828, 33656970,
				264907, 264908, 264909, 52644045,
				56182435, 265006, 265007, 60433438,
				55812038, 21906754, 52644296, 265010,
				264601, 265017, 265019, 264681, 264448,
			-	264682, 264288, 264686, 264687, 264688,
				21906766, 21906769, 55811957, 35695917.
				265020, 265021, 60170615, 264690, 264691,
				33657023, 264692, 264693, 65274620,
				27486264, 263972, 18108374, 18108377.
				264635, 264636, 264556, 60170394,
_				83373044, 65274727, 87168518, 22279000
2707	88089839 (5413, 5414)	88089839 (5413, 5414) Novel Protein sim. GBank gi]3417294 (AC004381) -		22278996, 22278998, 56182435, 21906754,
		Unknown gene product (Homo sapiens)		87168559, 265017, 264448, 52645129
2708		91011351 (5415, 5416) Novel Protein sim. GBank gij545790 bbs 147178 - DARPP-	UNCLASSIFIED	65274572, 264259, 29331822, 29331825.
		32-dopamine and cAMP-regulated phosphoprotein (human,		60432289, 29331826, 29331827, 29331828,
		brain, Peptide, 204 aa)		264909, 264510, 265007, 264910, 60433356,
				60433438, 33109954, 265010, 265011.
				264369, 264288, 264765, 264693, 264565
2709	94853988 (5417, 5418)	94853988 (5417, 5418) Novel Protein sim. GBank gij3169705 (AC004780) -	UNCLASSIFIED	29331822, 18108370, 18108374, 83373044
		F17127_1 [Homo sapiens]		
2710	87627979 (5419, 5420)			29331824, 264759, 264693, 18108382,
		(AL031432) dJ465N24.2.1 (PUTATIVE novel protein)		18108388
		(isoform 1) [Homo sapiens]		

gij3122400jspj035682IMUG_MOUSE - MYELOID UPREGULATED PROTEIN	Postsing professing demosis (DEONEGLE)	Iransforação	21906764, 21906765, 21906767, 21906768, 21906769, 55811957, 56994075, 22278997, 22278998, 2626789, 26278997, 26278997, 2627898, 264691, 264692, 33657023, 29331822, 29331824, 50424269, 29331826, 33657182, 26331824, 50424269, 29331826, 33657182, 263016, 264907, 60431850, 6043129, 264508, 264910, 60431850, 6043229, 264557, 26458, 55812038, 33109954, 18108385, 21906754, 33657084, 87168559, 60432113, 265017, 22279000, 265018, 264561, 18108351, 265018, 264766, 264766, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264788, 264768, 264768, 264768, 264788, 264768, 264768, 264768, 264768, 264768, 264
	Contains protein domain (PF00515) - Iransferase TPR Domain	iransierase	264486, 356962266, 222 78936, 264239, 29331824, 60432289, 35696052, 264508, 6043229, 35696052, 264506, 60432229, 33657402, 60433356, 265006, 265019, 18108351, 264681, 264288, 266685, 21906766, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 336571057, 35695917, 265020, 265021, 60170615, 264691, 264692, 33657023, 33657109, 33657182, 33657349, 18108370, 18108374, 35696423, 35695855, 264555, 52644332, 56182323, 87168518, 60432113
88003064 (5425, 5426) Novel Protein sim. GBank gi 2477513 (AC002398) - F25965_3 [Homo sapiens]		UNCLASSIFIED	66714117, 264906, 264563
13528218 (5427, 5428) 94122454 (5429, 5430) Novel Protein sim. GBank gil4321968 gb AAD15897  - (AF067430) Smarce1-related protein [Mus musculus]		UNCLASSIFIED	264636 264508, 264905, 264907, 264908, 264909, 264910, 264758, 265011, 264760, 264764, 264308, 364758, 364636
88003068 (5431, 5432) Novel Protein sim. GBank gil2477513 (AC002398) - F25965 3 [Homo sapiens]		glycoprotein	264091, 264259, 29331822, 66714117, 264908, 264369, 264693, 254556, 264563
80077461 (5433, 5434) Novel Protein sim. GBank gi[3327046 db] BAA31591  -   (AB014516) KIAA0616 protein [Homo sapiens]			264593, 264558
78604062 (5435, 5436)			264693
Novel Protein sim. GBank gil746495 (U23515) - weakly similar to gastrula zinc finger protein [Caenorhabditis elegans]		UNCLASSIFIED	29331822, 87168559, 265019, 265021, 52644150, 264691

2720	95086242 (5439, 5440)	2720   95086242 (5439, 5440) Novel Protein sim. GBank gil1335873 (U46690) - ATP-dependent RNA helicase [Mus musculus]	Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase		18108374, 60424179, 264469, 56182435, 21906765, 21906766, 35696423, 22278997, 265020, 265022, 265008, 265008, 264092, 264636, 60432229, 264691, 264692, 33657023, 264693, 33657402, 83373044, 29331824, 18108366, 60424269, 29331826, 37168474, 264100, 265010, 265011, 265019, 22373002, 264905, 264906, 18108351, 264681, 18108370, 29331830, 264908, 66712502, 2644045, 264909, 264909, 264908, 66712502, 2644045, 264909, 264909, 264908, 66712502, 2644045, 264909, 264909, 264908, 66712502, 2644045, 264909, 264909, 264909, 264908, 66712502, 2644045, 264909, 264828, 18108354
2721	95345523 (5441, 5442)	95345523 (5441, 5442) Novel Protein sim. GBank gil4929663lgblAAD34092.1 AF15185 - (AF151855) CGI-97 protein [Homo sapiens]	Contains protein domain (PF01172) - Uncharacterized protein family UPF0023		22278995, 35696286, 264259, 29331822, 29331824, 66714117, 29331826, 264906, 60433438, 265017, 18108351, 264448, 264288, 264769, 21906766, 265021, 33657109, 263969, 80431528, 264639, 55811576, 65274791, 35695655, 264631, 264637, 60170394, 56182323, 22279000
2722	91638807 (5443, 5444)	91638807 (5443, 5444) Novel Protein sim. GBank gil3212997[gblpAcC23434.1] - (AC004997) match to ESTs AA667999 (NID:g2626700), AA165465 (NID:g1741481), Z45871 (NID:g275105), and T84026 (NID:g712314); similar to various tre-like proteins including: AF040654 (PID:g2746883), D13644 (PID:g2104571), AL02114	Contains protein domain (PF00566) - oncogene TBC domain	oncogene	35696286, 22278999, 21906754, 265017, 264762, 264288, 21906765, 21906767, 21906768, 35695917, 18108362, 27486262, 35695855, 264558, 264559
2723	87387732 (5445, 5446)			UNCLASSIFIED	264508, 264509, 264906, 264909, 264910, 55812038, 264766, 264687, 264629, 264636, 264486
2724	87639563 (5447, 5448)	87639563 (5447, 5448) Novel Protein sim. GBank gil4680681lgblAAD27730.1JAF13295 - (AF132955) CGI-21 protein [Homo sapiens]		ubiquilin	18108396, 22278999, 20281099, 29331824, 29331826, 60432229, 60432289, 29331828, 60170831, 60432229, 60433438, 18108351, 264682, 21906766, 21906769, 35695917, 33657023, 33657109, 18108372, 18108374, 35695855, 22279000, 22279002
2725	94853991 (5449, 5450)	94853991 (5449, 5450) Novel Protein sim. GBank gi 3169705 (AC004780) - F17127_1 [Homo sapiens]		UNCLASSIFIED	264488, 52644507, 264259, 29331827, 21906754, 265011, 18108351, 264448, 264288, 264685, 264689, 35695917, 265020, 33657182, 27486261, 18108370, 18108374, 35696423, 18108385, 22279000
2726	86880599 (5451, 5452)	86880599 (5451, 5452) Novel Protein sim. GBank gil3342738 (AC005328) - R26660_1, partial CDS (Homo sapiens)		мнс	264468, 264828, 264685

### ### ### ### ### ### ### ### ### ##	727	91010470 (5453, 5454)	2727 91010470 (5453, 5454) Novel Protein sim. GBank gil731267[splP39219]RLUA_ECOLI - RIBOSOMAL LARGE SUBUNIT PSEUDOURIDINE SYNTHASE A (PSEUDOURIDYLATE SYNTHASE) (URACIL HYDROLYASE)	Contains protein domain (PF00849) - synthase RNA pseudouridylate synthase	synthase	62274572, 56182575, 22278994, 56994075, 22278999, 66432049, 29378996, 22278999, 64432049, 29331822, 29331824, 25278999, 66714117, 29331826, 29331827, 35696052, 29331828, 33656970, 264509, 66712502, 264910, 3365702, 264767, 21906764, 5581208, 21906764, 55811386, 265018, 265019, 264767, 21906765, 21906767, 21906769, 55811957, 35695917, 52644150, 33657023, 33657109, 33657109, 33657109, 33657109, 33657109, 264630, 65811876, 35696423, 35695855, 264630, 6043811876, 32279000, 26486, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 204286, 20428
Contains protein domain (PF00153) - UNCLASSIFIED Mitochondrial carrier proteins Contains protein domain (PF00753) - transport Mitochondrial carrier proteins  Contains protein domain (PF00795) - Carbon-nitrogen hydrolase Contains protein domain (PF00153) - UNCLASSIFIED  Mitochondrial carrier proteins  UNCLASSIFIED  Contains protein domain (PF00170) - transcriptfactor bZIP transcription factor	2728		Novel Protein sim. GBank gij3890433jembjCAA91399j - (Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene (Caenorhabdilis elegans)	Contains protein domain (PF00153) - I Mitochondrial carrier proteins	UNCLASSIFIED	181093, 25182435, 21906767, 55811957, 15585855, 265021, 264690, 264556, 264259, 264557, 29331822, 264559, 264448, 264288
94126026 (5459, 5460) Novel Protein sim. GBank gij3880433jemb CAA91399  - Contains protein domain (PF00153) - transport (266221) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene (26622) Novel Protein sim. GBank gij380433jemb CAA91399  - Contains protein domain (PF00155) - UNCLASSIFIED (266221) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene (266521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene (266524) Novel Protein sim. GBank gij4519821[dbj BAA75670.1] - Contains protein domain (PF00170) - transcriptitactor (AB017614) OASIS protein [Mus musculus]	729		Novel Protein sim. GBank gi[2408095 emb CAB16300  - (299168) putative RNA splicing protein  Schizosaccharomyces pombe}	Contains protein domain (PF00153) - I Mitochondrial carrier proteins	UNCLASSIFIED	65274572, 264259, 60432289, 66712502, 56182435, 264448, 264288, 264369, 55811957, 265021, 264557, 60432113
### Contains protein domain (PF00795) - ###################################	730	94126026 (5459, 5460)	Novel Protein sim. GBank gij3880433 emb CAA91399  - (Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabditis elegans]	Contains protein domain (PF00153) - Mitochondrial carrier proteins	transport	264887, 29331824, 29331826, 35696052, 264107, 56182435, 265008, 265009, 264592, 60431735, 265011, 264601, 265017, 18108351, 264288, 29148627, 55811957, 265021, 264690, 18108368, 18108387, 264558, 264559, 18108387, 56526486, 264566, 264486
94126028 (5463, 5464) Novel Protein sim. GBank gij3880433jemb CAA91399  - Contains protein domain (PF00153) - UNCLASSIFIED (266521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabditis elegans] 87363060 (5465, 5466) UNCLASSIFIED 94140286 (5467, 5468) Novel Protein sim. GBank gij4519621 dbj BAA75670.1  - Contains protein domain (PF00170) - transcriptfactor bZIP transcription factor	731		85	Contains protein domain (PF00795) - Carbon-nitrogen hydrolase		264259, 35696052, 265006, 264758, 264762, 264448, 264288, 29148627, 21906769, 87168518, 22279002
87363060 (5465, 5466) UNCLASSIFIED 94140286 (5467, 5468) Novel Protein sim. GBank gil4519621 dbj BAA75670.1  - Contains protein domain (PF00170) - Iranscripttactor (AB017614) OASIS protein [Mus musculus] bZIP transcription factor	732		Novel Protein sim. GBank gij3880433[emb]CAA91399] - (266521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabdilis elegans]	Contains protein domain (PF00153) - Mitochondrial carrier proteins	UNCLASSIFIED	264687, 264489, 18108358, 56182435, 264689, 35696423, 55811957, 265021, 265006, 265008, 264910, 265009, 264690, 264555, 264259, 264256, 264557, 264558, 264559, 18108383, 33657109, 87168518, 265010, 264601, 60432113, 265017, 264905, 264448, 263972, 264369, 264567
TOURDOO COUNTY CONTROL	734		Novel Protein sim. GBank gil4519621 dbj BAA75670.1  - (AB017614) OASIS protein [Mus musculus]	Contains protein domain (PF00170) - I bZiP transcription factor	UNCLASSIFIED	29331825, 264509, 264909 60424179, 52644507, 56182575, 264259, 29331828, 264907, 264510, 264910, 60433356, 265019, 55811150, 264681, 264763, 264681,

2	00131 00001660	1000				
	80+6) 00071 770	9, 94/U)	(2.33) (97716330 (3468), 3470) Novel Protein sim. GBank gij3830359 (ACU03278) - ESTS (95)		glycoprotein	22278996, 60432289, 29331827, 29146498, 264108, 264909, 264112, 33657402,
			gene. [Arabidopsis thaliana]	٠		87168474, 265017, 264762, 264448, 264764,
						264684, 21906765, 264693, 33657109,
						2009/0, 204030, 204030, 204037, 22279000, 1
2736	80247655 (5471, 5472)	1, 5472)			UNCLASSIFIED	264905, 264628, 264629, 263978, 264632,
$\neg$						264564
_	87604526 (5473, 5474)	5474)				264690
_	85731808 (5475,	5, 5476)	85/31808 (3475, 5476) Novel Protein sim. GBank gil2558501 dbj BAA22896  -  (D63850) hepatoma-derived growth factor [Mus musculus]			264488, 265009, 264768, 264691
2739	94319834 (5477,	7, 5478)	94319834 (5477, 5478) Novel Protein sim. GBank gi[5420387 emb CAB46679.1] -		UNCLASSIFIED	264684 83373044 264566
			(AJ243459) proteophosphoglycan (Leishmania major)			1000 (con cont)
2740	94148762 (5479,	9, 5480)	94148762 (5479, 5480) Novel Protein sim. GBank gij3417386 emb CAA75495		UNCLASSIFIED	264488, 56182575, 22278995, 35696286,
_			(Y15197) microtubule-associated protein, MAP-115 [Mus			22278997, 22278998, 22278999, 264259,
			muscalus			29331822, 29331824, 29331825, 29331827,
		_				35696052, 29331828, 29146498, 29331830,
						265006, 265007, 265009, 60432229,
	-	_				33657402, 55812038, 87168474, 265010,
_		_				265011, 265017, 265018, 265019, 264605,
_		_				264681, 264288, 264369, 52644229,
		_				21906765, 21906766, 21906767, 21906768,
		_				21906769, 265020, 265022, 264691, 264692,
						33657109, 18108370, 18108374, 55810764,
		_				35695855, 264634, 60431850, 264639,
		_				56182323, 18108382, 18108385, 65274727.
_	00047540	10073				22279002, 264564
14/7	8604/518 (5481,	. 5482)	8804/318 (3481, 3482) Novel Protein sim. GBank gij3242764 (AC005154) - similar		UNCLASSIFIED	22278996, 52644045, 52644229, 21906768,
_	,		61306) [Homo sapiens]			21906769, 265020, 60170615, 264691
2742	87648644 (5483,	3, 5484)			transferase	264259, 264905, 264758, 55812038, 264369,
			ALN - UDP-N-acetyl-alpha-			29148627
			U-galactosamine:polypeptide N- aretylpalactosaminylirans(arase 2 (CalNAc.T2)	beta-chain, 3 copies.		
2743	87627991 (5485	5486)	87627991 (5485, 5486) Novel Protein sim GRank gild468311lemblC4B370021.		LINICI ACCIETED	300350 300750 300750 367008 3600936
		•	(AL031432) dJ465N24.2.1 (PUTATIVE novel protein)			50050200, 201203, 201300, 201300, 200000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 20130000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 20130000, 2013000, 2013000, 2013000, 2013000, 2013000, 2013000, 20130000, 2013000, 2013000, 2013000, 2013000, 20130000, 20130000, 20130000, 20130000, 20130000, 20130000, 20130000, 20130000, 201300000, 20130000, 20130000, 20130000, 20130000, 20130000, 20130000000, 201300000, 20130000, 20130000, 201300000, 201300000, 20130000000000, 201300000000000000000000000000000000000
			(isoform 1) [Homo sapiens]			264764, 264288, 21906765, 21906767,
		_	•			264690, 264691, 264693, 263969, 263971,
						35695855 264637 264558 18108382
						60432113
2744	94126030 (5487,	, 5488)	94126030 (5487, 5488) Novel Protein sim. GBank gij3880433jemb CAA91399  -	Contains protein domain (PF00153) - kinase	kinase	18108374, 264488, 56182435, 21906765,
			(268521) similar to mitochondrial RNA splicing MSR4 like	Mitochondrial carrier proteins		35696423, 35695917, 35695855, 265020,
			protein; cDNA EST EMBL: C09217 comes from this gene			265021, 264511, 265009, 264490, 264556.
_			(Caenorhabditis elegans)			264259, 264557, 56182323, 264558, 264559,
_					•	18108383, 29331824, 18108385, 33657109,
_						29331826, 21906754, 29331827, 29331828,
						33657349, 87168518, 265018, 264905,
7						264482, 264448, 264486, 264369, 264288

2745	87740125 (5489, 5	5490)	2745 [87740125 (5489, 5480) Novel Protein sim, GBank gil4405795(gb)AAD19826(	Contains protein domain (PF00271) - helicase	helicase	35696286, 264509, 264905, 264907, 264908.
			(AF038963) RNA helicase [Homo saplens]	Helicases conserved C-terminal		264909, 264510, 264512, 265008, 264758,
				domain		264601, 265017, 264604, 264763, 264288,
						264686, 264769, 264693, 35696423,
						35695855, 264634, 264636, 264563, 264564.
						264565
2746		5492)	95418601 (5491, 5492) Novel Protein sim. GBank	Contains protein domain (PF00320) - UNCLASSIFIED	UNCLASSIFIED	22278996, 22278998, 22278999, 29331822,
		<u> </u>	gil4758738 ref NP_004680.1 pMTA1 - metastasis	GATA zinc finger		29331826, 29331827, 35696052, 29331828,
			associated 1			264905, 264906, 264907, 264908, 264909.
						52644045, 265006, 60170831, 264596,
						55812038, 265018, 264683, 264288,
						21906765, 21906767, 21906768, 21906769,
					, (·	265020, 264690, 33657023, 264693,
						33657109, 18108368, 18108374, 264558,
	_					18108385, 22279000, 264563
2747		5494)	94112677 (5493, 5494) Novel Protein sim. GBank		glycoprotein	264569, 52644507, 18108394, 22278995,
		<u> </u>	gil4557803 ref NP_000262.1 pNPC1 - Niemann-Pick			35696286, 22278997, 22278999, 52645080,
_		J	disease, type C1			29331624, 56182181, 29331826, 29331827,
						35696052, 264907, 264908, 264909, 265009,
						33109954, 55811386, 87168474, 265010,
						87168559, 264603, 265019, 264760, 264686,
						264768, 21906769, 35695917, 60170615,
						264692, 33657023, 52645129, 27486264,
						60431528, 18108374, 35696423, 35695855,
						264556, 56182323, 18108385, 264482
2748	91214983 (5495, 5	5496)	2748 [91214983 (5495, 5496) Novel Protein sim. GBank gil4191272 emblCAA099841 -	Contains protein domain (PF00646) -		65274572, 29331828, 264112, 264511,
		<u>~</u>	(AJ012295) apaG protein (Rhizobium etli)	F-box domain.		265019, 264760, 264767, 264768, 264769,
						21906768, 21906769, 265020, 27486262,
	,					56526486, 87168518, 22279000
2749	87346307 (5497, 5498)	5498)				264259, 264908, 264510, 265008, 265009.
						264760, 264369, 264768, 264563
2750		5500)	87336344 (5499, 5500) Novel Protein sim. GBank gij1872498 (U74297) - PiUS		UNCLASSIFIED	264488, 52644507, 18108396, 56994075.
		<u></u>	[Oryctolagus cuniculus]			264259, 29331825, 29331826, 29331827.
						29331828, 264508, 265009, 264910, 264591.
						264595, 33657084, 265011, 265019,
						18108351, 264288, 264686, 264769, 264689,
						55811957, 264693, 27486264, 18108370,
						18108374, 264558, 18108385, 264482,
				, 1		264563
2751		. 5502)			UNCLASSIFIED	29331822, 29331824, 265017, 33657023
2752		5504)	88062675 (5503, 5504) Novel Protein sim. GBank gij3041859 (AC004534) - OG-2	Contains protein domain (PF00046) - homeobox	homeobox	
		~	homeodomain protein-like; similar to U65067	Homeobox domain		
		1	(FID.9137332) Invitio Sapiens			

56182575, 56994075, 22278998, 29331822, 29331824, 29331825, 29331826, 265007, 264593, 55812038, 33109954, 18108351, 264288, 56181562, 21906767, 21906768, 265021, 264693, 18108374, 65274791, 264632, 56182323, 22279002, 264563, 264532, 264563, 264532, 264563, 264532, 264563, 264532, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264583, 264632, 264583, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 264632, 2646422, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264642, 264	22278993, 007 14 17, 2333 1027, 230330522, 29331828, 264906, 264908, 264909, 265011, 265017, 265019, 265019, 264288, 21906765, 21906767, 265022, 33657023, 264693, 56182323, 18108382, 22279000		264594	85658542, 264693	264693	65274572, 35696286, 66714117, 29331828, 264508, 56182435, 21906754, 55811957, 264629, 264636, 56182323, 22279002	264908	22278998, 29331622, 29331630, 265010, 265019, 265019, 264288, 21906765, 21906768, 21906769, 265020, 56182323, 22279002, 264563	18108394, 22278998, 264906, 264909, 265006, 265007, 264757, 265010, 265011, 265017, 265019, 18108351, 264448, 264683, 264686, 264768, 265020, 265021, 265022, 264691, 18108362, 264693, 18108365, 33657109, 18108368, 18108388, 87168518	264369, 35696423	52646842, 264259, 29331822, 29331825, 29331826, 29331826, 29331828, 33555970, 264905, 264907, 29331830, 265006, 265009, 21906754, 265019, 264448, 21906769, 27486262, 56182323, 56526486, 87168518, 264487
UNCLASSIFIED	Siruci	transcriptfactor	UNCLASSIFIED	eph -	struct		UNCLASSIFIED	nuclease	struct	- transcriptfactor	
Signature   Footalins protein domain (PF00646) - UNCLASSIFIED   Ovegicus   F-box domain.   F	535428 (U13736) - calmodulin- (Contains protein domain (PF00035) - struct EF hand	Contains protein domain (PF00096) - transcriptlactor Zinc finger, C2H2 type		Contains protein domain (PF00017) - leph Src homology domain 2				Contains protein domain (PF00023) - Inuclease Ank repeal	Contains protein domain (PF00989) - struct PAS domain	Contains protein domain (PF01352) - transcriptfactor KRAB box	Contains protein domain (PF00370) - FGGY family of carbohydrate kinases
94138972 (5505, 5506) Novel Protein sim. GBank gi 3851648 (AF098301) - neural F box protein NFB42 [Rattus norvegicus]	94115513 (5507, 5508) Novel Protein sim. GBank gi 535428 (U13736) - calmodulin- like protein [Pisum sativum]	)] Novel Protein sim. GBank gi[2996653 (AC004510) -  R30385_2 [Homo sapiens]	(6	95391590 (5513, 5514) Novel Protein sim. GBank gi[1173539 (U30473) - putative src-like adapter protein; non-catalytic src-like adapter protein containing SH3 and SH2 domains; homolog of mouse SLAP; Method: conceptual translation supplied by author [Homo sapiens]	79637846 (5515, 5516) Novel Protein sim. GBank gij3608372 (AF053768) - brain specific cortactin-binding protein CBP90 [Rattus norvegious]	91005312 (5517, 5518) Novel Protein sim. GBank gi 2072200 (U94863) - p40 [Borna disease virus]		87639597 (5521, 5522) Novel Protein sim. GBank gil4914573lemb[CAB43685.1  - (AL050390) hypothetical protein [Homo sapiens]	87592699 (5523, 5524) Novel Protein sim. GBank gij3136150 (AF050182) - PERIOD 3 [Mus musculus]	87539968 (5525, 5526) Novel Protein sim. GBank gij3511122 (AF060503) - zinc finner protein Homo sapiensi	94305140 (5527, 5528) Novel Protein sim. GBank gij2905643 (AF045244) - ribilol kinase [Klebsiella pneumoniae]
94138972 (5505, 5506)		88001472 (5509, 5510)	_				70824798 (5519 5520)				
2753	2754	2755	2756	2757	2758	2759	2780	2761	2762	2763	2764

2765	94315105 (5529, 5530)	2765   94315105 (5529, 5530) Novel Protein sim. GBank gil4688672 emb CAA17688.2			264488, 65274572, 22278995, 35696286,	Г
		/(match=(desc: fDrosophila melanopasier)			22278998, 22278999, 264259, 29331822,	
					29331824, 35696052, 264508, 264509,	
					2849US, 2649UB, 2649U7, 2649UB, 2649UB, 284543 284040 266000 264503 264503	
	-			-	204312, 204910, 203009, 204592, 204595, 264769 66812028 32100064 266040	_
					2047.30, 33012030, 33103934, 203010, 87168550 364600 385018 364750 364761	
					264762 264763 264448 264764 264789	-
					264766 264767 264768 56181562	
					21906764, 21906765, 21906768, 35695917.	
					265021, 264691, 264692, 33657023,	
					33657109, 264628, 18108370, 264629,	
					18108374, 55811576, 35696423, 35695855,	_
					264631, 264634, 264635, 264636, 264637,	_
					264638, 264558, 264639, 83373044,	_
					87168518, 22279000, 22279002, 264563,	
2768	94315109 (5531 5532)	Novel Destain aim Const all Edd 1011 - 1-100 Const			264482	
3			Contains protein domain (PF00076) - dna_ma_bind	dna_ma_bind	264488, 65274572, 60432289, 264907,	
		לייססססס וויסטורים והפון ליסטוסי ויססססס וויסטוסיוים ופוווויים ופון	RNA recognition motif. (a.x.a. KRM,		264909, 264511, 264512, 60433356, 264288,	_
			KBU, or KNP domain)		264685, 264689, 35695917, 265022, 264693,	_
					264628, 65274791, 264635, 264555, 264558,	_
					264557, 264638, 264558, 264559, 83373044,	_
27.67	RN204207 (5533 5534)	Octobrie de Contraction de Contracti			60432113	_
		cozovzer (1955), 2054) invoer Frotein sim. Gbank gil 10/9451 ipirijA55453 - tropomodulin, skeletal muscle - chicken		struct	264112, 263974, 264558	
2768		94322238 (5535, 5536) Novel Protein sim. GBank gi[5441322]emb[CAB46721.1] -		UNCLASSIFIED	264488, 263994, 264489, 35696286	Т
		((AL031427) dJ167A19.1 (novel protein) [Homo sapiens]			22278998 22278999 264259 29331822	
					60432289, 29331826, 35696052, 264508,	
					264509, 264905, 264906, 264907, 29331830,	
					264908, 264909, 264510, 264511, 264512,	
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					264681, 18108351, 264764, 264288, 264766,	
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					21906766, 21906768, 21906769, 35695917,	
				-	265021, 265022, 33657023, 33657109,	
					264628, 264629, 18108374, 35696423,	
					35695855, 264630, 264631, 264632, 264634,	
					264635, 264636, 264555, 264638, 264639,	
					83373044, 264563, 264564, 264565, 264566,	
					264486, 264567	٦

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tubulin	Kinase	UNCLASSIFIED	histone	UNCLASSIFIED	ribosomalprot
	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		Contains protein domain (PF00850) - Histone deacetylase family		Contains protein domain (PF00380) - inbosomalprot Ribosomal protein S9/S16
2769   95311088 (5537, 5538)   Novel Protein sim. GBank gi[5419859 emb CAB46375.1  - (AL096725) hypothetical protein [Homo sapiens]	87730182 (5539, 5540) Novel Protein sim. GBank gi 5701965 emb CAB52157.1  - (AL 109736) WD repeat prolein [Schizosaccharomyces pombe]	88084071 (5541, 5542) Novel Protein sim. GBank gil3093433 (AC004125) - Unknown gene product [Homo sapiens]	95357309 (5543, 5544) Novel Protein sim. GBank gil4885531  ref NP_005465.1 pNY Contains protein domain (PF00850) - histone C - histone deacetylase 5  Histone deacetylase family	94138994 (5545, 5546) Novel Protein sim. GBank gil3288888 (AC005253) - R26445_1 [Homo sapiens]	2774 87819908 (5547, 5548) Novel Protein sim. GBank gil465852(spjp34388)YLS3_CAEEL - HYPOTHETICAL 70.7 Ribosomal protein S9/S16 KD PROTEIN F09C8.3 IN CHROMOSOME III

2775		95307987 (5549, 5550) Novel Protein sim. GBank gil4689132[gb]AAD27775.1 AF07704 - (AF077042) 30S ribosomal protein S7 homolog [Homo sapiens]	Contains protein domain (PF00177) - ribosomalprot Ribosomal protein S7p/S5e	ribosomalprot	264488, 22278995, 56994075, 22278996, 35696286, 22278999, 264259, 29331822, 29331824, 29331825, 60432289, 29331827, 29331827, 29331827, 29331827, 29331827, 29331827, 265017, 21906754, 265019, 264449, 264682, 264369, 21906765, 21906766, 21906767, 21906768, 21906766, 21906767, 265022, 60170615, 264691, 18108370, 35696423, 65274791, 35695855, 264634, 60431850, 60170394, 56182323, 264558, 34486, 264657
2778	87791557 (5551, 5552)				56182575, 22278998, 22278999, 264259, 29331822, 29331824, 264908, 29331830, 264510, 33657402, 21906754, 55811386, 265017, 265019, 264448, 264289, 21906765, 21906766, 21906766, 21906766, 21906766, 258020, 265021, 265022, 60170615, 255020, 265201, 265052, 264555, 56526486, 257270000
1112	79818729 (5553, 5554)			UNCLASSIFIED	264907, 264766
2778				UNCLASSIFIED	264907, 264593, 264760, 264628
2779		87649729 (5557, 5558) Novel Protein sim. GBank gil4680711[gb]AAD27745.1 AF13297 - (AF132970) CGI-36 protein [Homo sapiens]		UNCLASSIFIED	22278997, 264259, 29331824, 66714117, 35696052, 265006, 264512, 264448, 264288 29148627, 18108364, 20281149, 18108370, 264629
2780		94679397 (5559, 5560) Novel Protein sim. GBank / gli4758524[ref]NP_004825.1[pHGK] - HPK/GCK-like kinase	Contains protein domain (PF00780) - kinase CNH domain	kinase	29146499, 65274791, 264634, 264639
2781		Novel Protein sim. GBank gil4469352 gb AAD21222  - (AF069502) ubiquitin specific protease UBP43 [Mus musculus]	Contains protein domain (PF00443) - t Ubiquitin carboxyl-terminal hydrolase family 2	ubiquitin	60424179, 29331824, 60424269, 66714117, 29331826, 56182435, 87168474, 265017, 264764, 56181562, 21906765, 21906768, 35695917, 265020, 33657023, 35695855, 56182323, 87168518
2782	94233146 (5563, 5564)	94233146 (5563, 5564) Novel Protein sim. GBank gil4505013 ref NP_002310.1 pLRN  - leucine-rich neuronal protein	Contains protein domain (PF00560) - struct Leucine Rich Repeat	struct	65274572, 22278996, 22278998, 60432049, 264259, 29331822, 29331824, 29331826, 60432289, 29331822, 29331824, 29331826, 60432289, 29331828, 264905, 264907, 265408, 264909, 26409, 26409, 26409, 26409, 26409, 26409, 26409, 26469, 21906768, 26499, 255043, 25504229, 264699, 21906768, 26499, 2355702, 264693, 263978, 3565789, 3355703, 264659, 263978, 35698855, 264557, 56182323, 83373044, 18108385

2783	80016629 (5565, 5566)			tm7	264909, 264628, 263978, 263981
2784					264259, 29331822, 29331824, 29331825, 264482
2785			,	ASSIFIED	22278999, 264908, 264768, 265018, 264769, 21906765, 21906768, 21906769, 265020, 264564
2786		<u>.</u>	2073564 (U80223) - eukaryotic Contains protein domain (PF00069) - kinase inase; DGCN2 (Drosophila Eukaryotic protein kinase domain		264905, 265017
2787	87901266 (5573, 5574)			UNCLASSIFIED	264768, 21906765, 21906768, 55811957, 22278999, 264093, 60170615, 264259, 29331822, 18108365, 29331824, 33657109, 29331827, 35696052, 264100, 264105, 264908, 263977, 55811576, 264635, 265017, 265019, 22279002, 55811150, 264369, 264288
2788		88090644 (5575, 5576) Novel Protein sim. GBank gij3252826 (AC004382) - Unknown gene product [Homo sapiens]		UNCLASSIFIED	
2789	85491275 (5577, 5578)	85491275 (5577, 5578) Novel Protein sim. GBank gij2495729jsp Q92556jY281_HUMAN - HYPOTHETICAL PROTEIN KIAA0281 (HA6725)		UNCLASSIFIED	264103, 21906769, 264693
	87602784 (5579, 5580)	Novel Protein sim. GBank gi 5101772 emb CAB45135.1  - (AJ242978) p621 [Homo sapiens]		struct	264488, 264769, 264910, 264631, 264636, 264690, 264691, 264259, 264638, 29331824, 264508, 264509, 264905, 264563, 264908, 264628, 18108370, 264907, 264764, 264908, 264288, 264909
2791		88083195 (5581, 5582) Novel Protein sim. GBank gi 2911266 (AC002550) -   Unknown gene product [Homo sapiens]			21906764, 18108368
2792		95083783 (5583, 5584) Novel Protein sim. GBank gil2854163 gb AAC02581.1  - (AF045642) No definition line found [Caenorhabditis elegans]			22278996, 22278997, 264259, 29331822, 29331824, 29331826, 66714117, 29331822, 66714117, 29331826, 66714262, 264906, 66712502, 29331837, 35696052, 264906, 66712502, 29331830, 264909, 60433256, 60433438, 3109954, 265010, 265011, 265017, 265018, 265019, 264682, 264011, 265017, 265018, 265020, 255021, 33657023, 263974, 18108374, 65274791, 35695855, 264556, 264558, 56182323, 83373044, 181083385, 56526486, 60432113, 22279000, 264567
2793	87425476 (5585, 5586)			UNCLASSIFIED	264259, 60432289, 66712502, 265009, 264636
2794	85794830 (5587, 5588)			UNCLASSIFIED	264689, 265022, 264691, 18108388, 264567

- ubiquitin 627/45/2, 501825/5, 35095200, 22278999, 60432049, 264259, 29331822, 29331825, 66714117, 60432289, 29331822, 29331825, 66714117, 60432289, 29331826, 29331827, 3669052, 29331828, 264907, 66712502, 56182435, 264511, 265007, 264512, 264910, 60432229, 60433356, 60433438, 33109954, 8558542, 265018, 265019, 26428, 264686, 21906764, 21906765, 21906768, 21906769, 25811957, 265020, 265021, 265020, 55811957, 265020, 265021, 264682, 65274791, 33657109, 35656486, 60432113, 22279002, 264823, 264567, 264687, 264563, 264564, 264567			UNCLASSIFIED 264488, 264490, 264259, 264448, 20281149, 20281152, 264556, 264557, 264558, 264559,
Contains protein domain (PF00179) - ubiquitui Ubiquitin-conjugating enzyme		Contains protein domain (PF00569) - UNCLASSIFIED Zinc finger present in dystrophin, CBP/p300	
Novel Protein sim. GBank gi[5454146 ref NP_006348.1 pUBE2 - ubiquilin-conjugating enzyme E2E 3 (homologous to yeast UBC4/5)	94648857 (5591, 5592) Novel Protein sim. GBank gil4680651[gb]AAD27715.1[AF13294 - (AF132940) CGI-06 protein [Homo sapiens]	95110790 (5593, 5584) Novel Protein sim. GBank gil4838557lgblAAD31040.11 - (AF143859) potassium channel modulatory factor DEBT-91 [Mus musculus]	86198005 (5595, 5599) Novel Protein sim. GBank gil2852645 (AF007160) - unknown [Homo sapiens]
95334888 (5589, 5590)	2786 94848857 (5591, 5592) P	2797 95110790 (5593, 5594)	2798 86198005 (5595, 5596)

22278996, 22278997, 22278998, 22278999, 60432049, 264259, 29331826, 35696052, 29331828, 264909, 60433356, 33657402, 33109954, 87168474, 264448, 52644229, 21906766, 21906767, 21906768, 35695917, 265020, 265021, 265022, 52644150, 35695855, 264634, 60432113, 22279000	264488, 56994075, 264259, 20281099, 29331825, 29331827, 264905, 56182435, 265006, 265011, 87168559, 265017, 265019, 26448, 26428, 264766, 264686, 60170815, 264691, 264692, 27486265, 264628, 264629, 264636, 264562, 264562, 264564, 264566, 264567, 264567, 264568, 264567, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264688,	265007, 264687	264448, 35695855	264639	264566	264092, 264259, 29331822, 29331824, 29331827, 29331828, 264508, 265007, 265009, 21906754, 264369, 264288, 264632, 60170394, 264563, 264482	22278995, 22278996, 22278999, 264259, 29331822, 29331822, 29331826, 29331827, 29146498, 265008, 265009, 60433438, 265017, 265018, 265019, 264448, 264288, 21906765, 21906767, 21906769, 29148629, 265022, 52644150, 56182323
UNCLASSIFIED	glycoprotein		transport	UNCLASSIFIED	peplidase		UNCLASSIFIED
		Contains protein domain (PF00627) - UBA domain	ij2337865 (AC002464) - organic Contains protein domain (PF00083) - transport ilarity to JC4884 (PID:g2143892) Sugar (and other) transporter		Contains protein domain (PF01585) - peptidase G-patch domain		
2789   88090651 (5597, 5588)   Novel Protein sim. GBank gil3252825 (AC004382) - Unknown gene product [Homo sapiens]	88316481 (5599, 5600) Novel Protein sim. GBank gil4240301 dbj BAA74929.1  - (AB020713) KIAA0906 protein [Homo sapiens]	11, 5602)	B8082477 (5603, 5604) Novel Protein sim. GBank gi 2337865 (AC002464) - organic   Contains protein domain (PFC cation transporter; 50% similarity to JC4884 (PID:92143892) Sugar (and other) transporter   [Homo sapiens]	-	2804 57111131 (5607, 5608) Novel Protein sim. GBank gil4559368lgblAAD23029.1lAC00658 - (AC006585) hypothetical protein [Arabidopsis thaliana]		2806   87898951 (5611, 5612) Novel Protein sim. GBank , gij1168973jsp P44403jCLPB_HAEIN - CLPB PROTEIN
2799   88090651 (5597	2800 88316481 (5599	2801 86068814 (5601, 5602)	2802 88082477 (5603	2803 79577446 (5605, 5606)	2804 57111131 (560)	2805 87398486 (5609, 5610)	2806 87898951 (561:

ar ar 1988 Contains protein domain (PF00170) - bZIP transcription factor	UNCLASSIFIED 52644507, 52645156, 52646842, 18108398, 56182575, 22278994, 22278995, 22278996, 56182575, 22278994, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 264269, 529331822, 29331824, 29331825, 29331822, 29331826, 3365970, 264100, 264105, 264907, 5264045, 60433356, 264594, 60433438, 52646317, 2190674, 33057084, 52644296, 8168474, 265011, 87168559, 265017, 265018, 265019, 18108351, 264448, 264763, 264687, 52644269, 21906765, 21906766, 21906767, 21906768, 21906766, 21906767, 21906768, 269626, 27486262, 27486261, 27486262, 27486265, 35695763, 18108370, 18108377, 35696423, 35695855, 52644332, 83373044, 18108387, 87168518, 6143213	60432289, 264510, 265010, 265017, 265018, 264681, 264686, 265021, 264690, 22279000, 264586	ATPase_associated 18108351	UNCLASSIFIED 264569, 22278996, 264091, 264259, 29331828, 29146499, 29148629, 29148784, 264693			UNCLASSIFIED 56182575, 29331828, 264906, 66712502, 55811386, 265017, 265018, 265019, 264683, 265020, 87158518, 60473113		SSIFIED 264907, 264909	
ot 7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	CNC	MHC	ATPase	UNCLA	UNCLA	UNCLA	UNCLA	00170) - transcrip	UNCLA	CNCLA
Novel Protein sim. GBank gij4468310jembjCAB379911- [AL031432] dJ465N24.1 (PUTATIVE novel protein similar to predicted yeast and worm proteins) [Homo sapiens]  Novel Protein sim. GBank gij5541863jembjCAB51071.1]- [AL096857) hypothetical protein [Homo sapiens]  Novel Protein sim. GBank gij3254583 (AC005189) - match to ciliary dynein beta heavy chain; 78% Similarity to P23098 (PID:g118965) [Homo sapiens]  Novel Protein sim. GBank gij3264583 (AC005189) - match to ESTs H9778 (NID:g118965)] [Homo sapiens]  Novel Protein sim. GBank gij3264583 (AC005620) - R33590_1 [Homo sapiens]  Novel Protein sim. GBank gij3548791 (AC005620) - R33590_1 [Homo sapiens]  Novel Protein sim. GBank gij5420389jembjCAB46680.1] - (AJ243460) proteophosphogiycan [Leishmania major]								Contains protein domain (PFI		
91720702 (5613, 5614) 95359111 (5615, 5616) 88083530 (5617, 5618) 87259032 (5619, 5620) 97235845 (5621, 5622) 88093334 (5623, 5626) 91218755 (5625, 5626) 90980906 (5627, 5628) 95358229 (5631, 5632)	2807 91720702 (5613, 5614) Novel Protein sim. GBank gij4468310jembjCAB37991j - (AL031432) dJ465N24,1 (PUTATIVE novel protein similar to predicted yeast and worm proteins) [Homo sapiens]	Novel Protein sim. GBank gij5541863 emb CAB51071.1  - (AL096857) hypothetical protein [Homo sapiens]				Novel Protein sim. GBank gij3264583 (AC005189) - match to ESTs H97758 (NID:g1118643) and AA085546 [(NID:g1628773) [Homo sapiens]	Novel Protein sim. GBank gil4240273 dbj BAA74915.1  - (AB020699) KIAA0892 protein [Homo sapiens]	Novel Protein sim. GBank gij3548791 (AC005620) - R33590_1 [Homo sapiens]		Novel Protein sim. GBank gil5420389 emb CAB46680.1  - (AJ243460) proteophosphoglycan [Leishmania major]
	2807	2808	2809	2810	2811		2813	2814	2815	2816

		UNCLASSIFIED 264909, 264511		ment 264636	UNCLASSIFIED 264259, 29331822, 60432289, 264905, 60433356, 33657402, 265011, 264760, 21906765, 60170615, 264638	ment 264766	UNCLASSIFIED 264907	264760	52645156, 22278994, 35696286, 22278997, 22278998, 52645080, 29331822, 29331824, 29331825, 29331827, 35696052, 29331824, 3365970, 52644045, 52646317, 33657084, 52644296, 265017, 265018, 265019, 284288, 21906764, 21906765, 21906766, 21906767, 21906769, 35695917, 52644150, 33657109, 33657182, 27486261, 27486261, 3569565, 2564432, 3569565, 2564432, 3569565, 2564432, 3659565, 2564432, 3659565, 2564432, 3659565, 2564432, 3659565, 2564432, 3659565, 2564432, 36595621, 36596421, 36595626, 36595621, 36596421, 3659565, 2564432
	- UNCLA	UNCLA	- UNCLA	) - complei	UNCLA	- compler	UNCLA		
	Contains protein domain (PF00023) - UNCLASSIFIED Ank repeat		Contains protein domain (PF00919) - UNCLASSIFIED Uncharacterized protein family UPF0004	Contains protein domain (PF00386) - complement C1q domain		Contains protein domain (PF00386) - complement C1q domain			
2817   87749542 (5633, 5634) Novel Protein sim. GBank gi[1293846 (U56966) - coded for by C. elegans cDNA yk30b3.5; coded for by C. elegans cDNA yk30b3.3 [Caenorhabditis elegans]	88073579 (5635, 5636) Novel Protein sim. GBank gil549986 (U13149) - possible apospory-associated protein [Pennisetum ciliare]		2820   87765744 (5639, 5640) Novel Protein sim. GBank gil4929773 gb AAD34147.1 AF15209 - (AF152097) CGI-05 protein [Homo sapiens]	95320511 (5641, 5642) Novel Protein sim. GBank gij399144 sp P02747 C1QC_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, C CHAIN PRECURSOR	94260221 (5643, 5644) Novel Protein sim. GBank gi[2224671 dbj BAA20820  - (AB002363) KIAA0365 [Homo sapiens]	95320513 (5645, 5646) Novel Protein sim. GBank gij399144 sp P02747 C1QC_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, C CHAIN PRECURSOR	95320515 (5647, 5648) Novel Protein sim. GBank gij399144 sp P02747 C1QC_HUMAN - COMPLEMENT C1Q SUBCOMPONENT, C CHAIN PRECURSOR		94311905 (5651, 5652) Novei Protein sim. GBank gil3859683 emb CAA22020  - (AL033503) conserved hypothetical protein [Candida afbicans]
37749542 (5633, 5634)	38073579 (5635, 5636)	87793527 (5637, 5638)	87765744 (5639, 5640)	95320511 (5641, 5642)	34260221 (5643, 5644)	35320513 (5645, 5646)	95320515 (5647, 5648)	19742170 (5649, 5650)	94311905 (5651, 5652)
2817 (	2818	2819 8	2820	2821 9	2822   8	2823 9	2824 9	2825	2826

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95320519 (5653, 5654) Novel Protein sim. GBank gil tyrosine phosphatase TD14 [ 91229615 (5655, 5656) Novel Protein sim. GBank gil tyrosine phosphatase TD14 [ 91229615 (5655, 5656) Novel Protein sim. GBank gil tyrosine phosphatase TD14 [ 91229615 (5657, 5658) Novel Protein sim. GBank gil tyrosine [Homo sapiens] 91249667[sp]G61200[NPH1	Contains protein domain (PF00386) -	264486, 18108391   264486, 18108391   2654986, 18108391   2593974 (AF077000) - protein Contains protein domain (PF00102) - phosphatase   29331822, 35696052, 264104, 265409, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019, 265019,			UNCLASSIFIED 265017 UNCLASSIFIED 22278997, 22278999, 52646317, 264288, 264688, 21906767, 60431528, 264638, 22279000	11 - UNCLASSIFIED	dbj BAA76788.1  - ATPase_associated 60432289, 29331828, 265000, 265010, 265011, 264448, 55811957, 265020.   18108370
	ا ر'≨		2829 87651244 (5657, 5658) Novel Protein sim. GBank gij4680689jgb AAD27734.1 AF13295 protein [Homo sapiens]	88087109 (5659, 5660) Novel Protein sim. GBank gij2498667 sp Q61200 NPH1	2831 87614717 (5661, 5662) 2832 87631809 (5663, 5664)	2833 87612938 (5665, 5666) Novel Protein sim. GBank gi 5262615  (AL080156) hypothetical protein [Hom 2834 86974703 (5667, 5668) Novel Protein sim. GBank gi 2224567 (ABAND313 Horne sanion)	2835 87775712 (5669, 5670) Novel Protein sim. GBank gil4589532 (AB023161) KIA40944 protein [Homo

		UNCLASSIFIED 264555	264509, 264511, 265011, 264288, 264769, 265020, 264634, 264636, 264556	dna_rna_bind 56182575, 35696052, 264907, 264908, 264909, 264909, 264909, 264593, 264595, 264766, 265022, 264691, 33657182, 35695763, 18108370, 35695855, 264631, 264559, 264563, 264567	264600	UNCLASSIFIED 264906, 264907, 264908, 264910, 264764, 35695855, 83373044, 18108385	264685		UNCLASSIFIED 29331830, 264909, 265008, 265011, 87168559, 264629, 264556	UNCLASSIFIED 264908, 264766, 264769, 264629, 264637, 264566	UNCLASSIFIED 264490, 264259, 264508, 264905, 264907, 264510, 265007, 265008, 264591, 264592, 264593, 264594, 264595, 55812038, 265010, 265011, 264604, 264763, 264764, 264765, 264766, 264628, 264629, 264556, 264556, 264559, 264559, 264559, 264559, 264559, 264569, 264559, 264567		UNCLASSIFIED (35696286, 265008, 265009, 265018, 264288,   35695817, 264693, 18108374, 35695855.
		IUNO		Contains protein domain (PF00076) - dna. RNA recognition motif. (a.k.a. RRM., RBD. or RNP domain)		ONO.	Contains protein domain (PF00008) - EGF-like domain	нон	אס	ONO	חאכ		n
2837	87775392 (5675, 5676) Novel Protein sim. GBank gil973378 (U31263) - core protein [Hepatitis C virus]		87774665 (5679, 5680) Novel Protein sim. GBank gil1575515 (U64899) - thrombospondin-related anonymous protein [Plasmodium gallinaceum]	86982568 (5681, 5682) Novel Protein sim. GBank gi[2224605 dbj BAA20790  - (AB002330) KIAA0332 [Homo sapiens]		91012494 (5685, 5686) Novel Protein sim. GBank gi[5578957 emb CAB51350.1  - (AL050306) dJ475B7.1 (novel KIAA0615 and KIAA0323 LIKE protein) [Homo sapiens]	56731154 (5687, 5688) Novel Protein sim. GBank gij585123 sp Q08878 FBLC_MOUSE - FIBULIN-1, ISOFORM C PRECURSOR (BASEMENT-MEMBRANE PROTEIN 90) (BM-90)	94321719 (5689, 5690) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]	88318613 (5691, 5692) Novel Protein sim. GBank gij5306263[gbjAAD41995.1]AC00623 - (AC006233)	81811757 (5693, 5694) Novel Protein sim. GBank gij3399576 (AC005390) - R31180 1 [Homo sapiens]	87612943 (5695, 5696) Novel Protein sim. GBank gil5262615 emb CAB45747.1  - (AL080156) hypothetical protein [Homo sapiens]	88084283 (5697, 5688) Novel Protein sim. GBank gij3342218 (AC004131) - Unknown gene product [Homo sapiens]	
37 87766482 (5673, 5674)		2839 85799317 (5677, 5678)	87774665 (5679, 5680)		42 80080086 (5683, 5684)	2843 91012494 (5685, 5686)	2844 56731154 (5687, 5688)	2845 94321719 (5689, 5690)	2846 88318613 (5691, 5692)	2847 81811757 (5693, 5694)	2848 87612943 (5695, 5696)	2849 88084283 (5697, 5698)	2850 87623636 (5699, 5700)

2851	87820548 (5701 5702)	2851   87820548 (5701 5702)  Novel Protein sim CBank all 1234540  Protein sim			
		(AF051098) seven transmembrane domain orphan receptor [Mus musculus]		UNCLASSIFIED	264906, 264907, 56182435, 264758, 55811386, 265010, 18108351, 264448, 264369, 21906765, 264691, 264692, 264693,
2852	86987023 (5703, 5704)	86987023 (5703, 5704) Novel Protein sim. GBank gil1825729 (U88308) - similar to drosophila membrane protein PATCHED SP:P18502 (PID:0129645) (Caenorhabdiils elegans)		UNCLASSIFIED	3569555, 264556, 18108385, 264567 264591
2853	87784630 (5705, 5706)	2853 87784630 (5705, 5706) Novel Protein sim. GBank gi[2702347 (AF027503) - putative membrane-associated guanylate kinase 1 [Mus musculus]	Contains protein domain (PF00397) - kinase	kinase	56182575, 55811150, 264690, 27486262, 27486265, 264632, 56182323, 56526486.
2854		88083557 (5707, 5708) Novel Protein sim. GBank gi[2795825 (AC004021) - kelch protein; ring canal component involved in cytoplasmic bridges; 77% Similarity to A45773 (PID:g1079096) [Homo	Contains protein domain (PF01344) - dna_ma_bind Kelch motif	dna_ma_bind	22279002 35696286, 29331824, 29331826, 29331828, 264908, 264768, 264693, 22279002, 264482
2855		94723856 (5709, 5710) Novel Protein sim. GBank gi 1504040 db  BAA13219  - (D86983) similar to D.melanogaster peroxidasin(U11052) [Homo sapiens]	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	glycoprotein	2278994, 29331822, 29331824, 29331825, 264906, 264908, 33109954, 265018, 265019, 264448, 21906765, 265020, 264690, 27486265, 83373044, 22279000, 22279002,
2856		88093359 (5711, 5712) Novel Protein sim. GBank gij3264583 (AC005189) - match to ESTs H97758 (NID:g1118643) and AA085546 (NID:g162873) IHomo sapiens)			264482 21906766, 22278997, 265022, 29331822, 29331826, 27486262, 265007, 265009,
	95348286 (5713, 5714)	2 S S	3041855 (AC004537) - similar Contains protein domain (PF00628) - struct FHD-finger ins	siruct	205017, 20482, 264563, 18108351 22278995, 35696286, 29331824, 29331825, 35696052, 264103, 264108, 56182435, 21906765, 21906769, 265020, 18108368,
	87434748 (5715, 5716)		Contains protein domain (PF00097) - dna_rna_bind Zinc finger, C3HC4 type (RING finger)	dna_rna_bind	264569, 264867, 22278995, 22278996, 264569, 264867, 22278995, 22278996, 22278996, 22278996, 22278996, 22278996, 22278996, 22278996, 22278996, 22331827, 22331827, 23331826, 264509, 264509, 264905, 264906, 29331830, 264908, 52644045, 264900, 265011, 265001, 265001, 265001, 265011, 264004, 265011, 265001, 265011, 264004, 265018, 264766, 18108351, 264769, 264765, 264288, 264369, 264685, 264766, 264765, 264691, 265031, 264694, 264766, 264691, 265031, 264694, 264766, 264691, 265031, 264634, 264691, 18108370, 264692, 18108370, 264639, 18108381, 264631, 264596, 26279002, 264566, 264566, 264488, 264567
6007	90937675 (5717, 5718)	9093/6/5 (5/17, 5/18) Novel Protein sim. GBank gil4325320 gb AAD17331.1  - (AF124427) claudin-15 [Mus musculus]	1	UNCLASSIFIED	60424179, 65274572, 29331828, 264905, 264511, 264758, 265011, 21906767, 21906769, 55811957, 265021, 56182323

264259, 29331822, 29331824, 29331825, 29331826, 29331827, 29331828, 264510, 264511, 33109954, 18108351, 264683, 264765, 264369, 264686, 21906765, 264691, 264692, 264693, 18108388, 22279002,	264369, 264692	264691, 264638	56994075, 35696286, 22278998, 28331822, 29331824, 35696052, 29331828, 264106, 264511, 55812038, 33657084, 55811386, 265018, 265019, 21906765, 21906766, 21906769, 35695917, 265020, 265022, 33657023, 33657109, 33657349, 264629, 18108376, 60431850, 56182323, 18108385, 18108387, 87168518, 22279902	264259, 264910	22278995, 21906764, 264482	65274572, 22278996, 22278998, 22278999, 264259, 29331827, 29331828, 260433438, 21906754, 265018, 264448, 264764, 52644150, 83373044	265022, 264259, 264508, 22578998, 265022, 264259, 264508, 264907, 264511, 264910, 264635, 264636, 264637, 265011, 265017, 265019, 265019, 264563, 264088, 264566, 264764, 264369, 264567, 264486, 264288, 264766
UNCLASSIFIED		UNCLASSIFIED	siruci	struct		glycoprolein	UNCLASSIFIED
			Contains protein domain (PF00096) - struct Zinc finger, C2H2 type				
2860 87532599 (5719, 5720) Novel Protein sim. GBank gil4469186 emb CAB38414.1 - (AL031588) d.J1163J1.2.1 (novel protein similar to C. elegans B0035.16 and bacterial IRNA (5- Methylaminomethyl-2-thiounidylate)-Methyltransferases) ((soform 1) [Homo sapiens]	86698507 (5721, 5722) Novel Protein sim. GBank gil3941730 (AF108083) - BS4 [Homo sapiens]	2862   87569585 (5723, 5724) Novel Protein sim. GBank gil4505013 ref NP_002310.1 pLRN  - leucine-rich neuronal protein	6) Novel Protein sim. GBank gij3249088 (AC004473) - Contains sImilarity to goliath protein gb M97204 from D. metanogster. [Arabidopsis thaliana]	8/42/030 (5/27, 5/28) Novel Protein sim. GBank gi 10/9451 pir  A55463 - tropomodulin, skeletal muscle - chicken	95312191 (5/29, 5/30) Novel Protein sim. GBank gil438840 (L19048) - MSA-2 [Plasmodium falciparum]	95105480 (5731, 5732) Novel Protein sim. GBank gij585703jspjQ07066jPMP2_RAT - 22 KD PEROXISOMAL MEMBRANE PROTEIN	86908001 (5733, 5734) Novel Protein sim. GBank gil4580997[gb]AAD24571.1[AF12108 - (AF121081) CAMP inducible 2 protein [Mus musculus]
87532589 (5719, 572	86698507 (5721, 5722	87569585 (5723, 5724	91220421 (5725, 5726		95312191 (5729, 5730	- 1	
286(	2861	2862	286;	2864	CQR7	2866	2867

(X94232) t-Cell activation g 2869 88094412 (5737, 5738) 2870 84404574 (5739, 5740) 2871 88318621 (5741, 5742) Novel Protein sim. GBank gil5306263lgblAAD41995.	(X94232) t-Cell activation protein [Homo sapiens]	·	22278998, 22278999, 60432049, 264259, 29331822, 29331824, 29331825, 6043289, 29331826, 29331827, 35696052, 29331828, 264508, 52644045, 264828, 265006, 265007
9 88094412 (5737, 5 0 84404574 (5739, 5 11 88318621 (5741, 5			29331822, 29331824, 29331825, 60432289, 29331826, 29331827, 35996052, 29331828, 264808, 52844045, 264828, 265006, 265007
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			29331826, 29331827, 35696052, 29331828, 264508, 52644045, 264828, 265006, 265007
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5		·	264508, 52644045, 264828, 265006, 265007,
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			265008, 265009, 264591, 60432229, 264593.
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			60433356, 264595, 21906754, 265017,
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			265019, 264682, 264369, 21906765,
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			21906766, 21906767, 21906768, 265021,
59 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			265022, 264691, 33657182, 18108368,
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			27486261, 27486262, 27486264, 27486265,
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			18108370, 18108374, 35696423, 35695855,
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			264632, 56182323, 87168518, 264404,
69 88094412 (5737, 5 70 84404574 (5739, 5 71 88318621 (5741, 5			22279000, 22279002, 264482, 264563,
69 88094412 (5737, 5: 70 84404574 (5739, 5: 71 88318621 (5741, 5:			264564, 264567, 264487
70 84404574 (5739, 5 71 88318621 (5741, 5	738)	UNCLASSIFIED	264369
71 88318621 (5741, 5	740)	UNCLASSIFIED	264905, 264908, 264764, 21906769, 264634
	742) Novel Protein sim. GBank	UNCLASSIFIED	264259, 29331822, 60432289, 29331827.
	gi 5306263 gb AAD41995.1 AC00623 - (AC006233)		264907, 265008, 265017, 265018, 264682,
	unknown protein [Arabidopsis thaliana]		264764, 18108354, 265021, 27486265.
			264629, 18108387, 264567
2872   95312197 (5743, 5	95312197 (5743, 5744) Novet Protein sim. GBank gil112205 pir  B39066 - proline- rich protein 15 - rat	kinase	263981
2873 88094252 (5745, 5746)	746)	UNCLASSIFIED	264488, 18108374, 264768, 264687, 264688,
			264689, 35696423, 35696286, 35695917,
			264510, 264511, 265007, 264512, 265008.
			264910, 264534, 264634, 264635, 264555,
			264592, 264259, 264558, 60433438,
_	-		60432289, 35696052, 265011, 264600,
			264601, 60432113, 264508, 264563, 264482,
			264509, 264905, 264906, 264564, 18108351.
			264763, 18108370, 264907, 264566, 264908.
			264764, 264288, 264567, 264909, 264486,
			264766, 18108391

3212854 (AC004005) - s thaliana] 2781386 (AC004010) - similar   Contains protein domain (PF00560) - glycoprotein	ch Repeat	Contains protein domain (PF00167) - fgf Fibroblast growth factor		Contains protein domain (PF01207) - UNCLASSIFIED Uncharacterized protein family UPF0034
AC004005) -  AC004010) - similar	ch Repeat	domain (PF00167) - factor		in (PF01207) - n family
ana] ana] 386 (ACOD4010) - similar	Contains protein dom	Contains protein domair Fibroblast growth factor		Contains protein domain (PF0) Uncharacterized protein family UPF0034
2874 94313549 (5747, 5748) Novel Protein sim. GBank gij2212854 ( unknown protein [Arabidopsis thaliana] unknown protein [Arabidopsis thaliana]	Nover Protein sim. Gbank gijz/81/85 (ACO44010) - similar to Leucine-rich transmembrane proteins; 44% similarity to U42767 (PID:g1736918) [Homo sapiens]	88090854 (5751, 5752) Novel Protein sim. GBank gi 2979530 (AC004449) -  R33683_2 [Homo sapiens]	94747029 (5753, 5754) Novel Protein sim. GBank gil4704208 emb CAB41646.1  - (AL035419) dJ1100H13.1 (putative novel protein) [Homo sapiens]	88095309 (5755, 5756) Novel Protein sim. GBank gi[3876775 emb CAB03067  - (Z81077) predicted using Genefinder; Similarity to Yeast protein 8248 (TR:G587531) [Caenorhabditis elegans]
2874 94313549 (5747, 5749) N	86063 / Zo (3/49, 3/30)	2876  88090854 (5751, 5752)	94747029 (5753, 5754) ,	2878   88095309 (5755, 5756)

18108359, 264259, 264905, 18108370, 264629, 264609, 18108374, 18108377, 265006, 264910, 264637, 60170394, 264569, 265017, 264564, 264565, 264567, 264567, 264684, 264369	264488, 52646365, 52846842, 22278994, 35698286, 22278998, 22278999, 264259, 29331822, 29331826, 29331826, 29331826, 29331826, 29331827, 35696052, 29331828, 284107, 264508, 264509, 264909, 264909, 2644045, 264510, 265006, 264511, 265007, 264512, 265009, 264900, 264694, 21900754, 52846317, 25849296, 264764, 264569, 264600, 264768, 264769, 264769, 264769, 264769, 264769, 264697, 264769, 264692, 33657023, 264634, 264635, 264636, 264635, 264634, 264635, 264636, 264635, 264636, 264638, 264638, 264638, 264638, 264638, 264658, 264659, 264483, 264563, 264569, 264563, 264659, 264483, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264483, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264488, 264563, 264563, 264563, 264488, 264483, 264567, 264488	22278998, 29331822, 52644045, 21906765, 264639, 60432113	264488, 22278996, 22278999, 29331822, 29331826, 264908, 60170831, 60433356, 55812038, 264681, 264682, 264686, 264687, 264688, 21906769, 264693, 263967, 18108374, 55811576, 56182323, 22279002, 264566	264112, 264682	265018, 264634	264686, 264693	60432049, 264259, 29331828, 264908, 264511, 264595, 60433438, 264596, 265017, 264605, 263969, 263972, 264555, 83373044, 87168518, 264566	18108351, 264686, 264629, 264631, 264639, 83373044, 264482	264488, 264259, 29331828, 264508, 264906, 264593, 284593, 284758, 264766, 264769, 18108374, 83373044, 264486
UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED	complementrecept	UNCLASSIFIED	transcriptfactor
Contains protein domain (PF00315) - UNCLASSIFIED Uracil-DNA glycosylase				Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat			Contains protein domain (PF00084) - complementrecept Sushi domain (SCR repeat)		Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type
gil4895145[gb AAD32752.1  - eptomyces lavendulae]	94851439 (5759, 5760) Novel Protein sim. GBank gil4680703[gb AAD27741.1[AF13296 - (AF132966) CGI-32 protein [Homo sapiens]	87650539 (5761, 5762) Novel Protein sim. GBank gil733571 (U23452) - No definition line found [Caenorhabditis elegans]	87714367 (5763, 5764) Novel Protein sim. GBank gi 1118112 (U41559) - No definition line found [Caenorhabditis elegans]	95362875 (5765, 5766) Novel Protein sim. GBank gil4868008[gb]AAD31087.1 AF10693 - (AF106934) vitamin D receptor-interacting protein [Homo sapiens]			91237823 (5771, 5772) Novel Protein sim. GBank gil1255889 (U53344) - T07H6.5 gene product [Caenorhabditis elegans]	91227860 (5773, 5774) Novel Protein sim. GBank gil3882323 db  BAA34521.1  - (AB018344) KIAA0801 protein [Homo saplens]	95105816 (5775, 5776) Novel Protein sim. GBank gil4508015 ref NP_003447.1 pZNF2 - zinc finger protein 205 Zinc finger, C2H2 type
87869122 (5757, 5758)     (			87714367 (5763, 5764) i		87784643 (5767, 5768)	83006306 (5769, 5770)   	91237823 (5771, 5772)   {	91227860 (5773, 5774)	95105816 (5775, 5776) I
2879		2881	2882	2883	2884	2885	2886	2887	2888

0000	10750 6537 6330			COLUMN TO STATE OF THE PERSON NAMED IN COLUMN TO STATE OF	CC004076 20004004 200000 204700
	_				18108354, 264288
2890	78703853 (5779, 5780)	78703653 (5779, 5780) Novel Protein sim. GBank gil854065 emb CAA58337  -  (X83413) U88  Human herpesvirus 6		UNCLASSIFIED	264591, 264766
2891		88094428 (5781, 5782) Novel Protein sim. GBank gij3877750jembjCAB01508j - (Z78064) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D65865 comes from this gene; cDNA EST EMBL:D69451 comes from this gene; cDNA EST EMBL:D66026 comes from this gene; cDNA EST EMBL:D69658 comes from this gene; cDNA EST		UNCLASSIFIED	264591, 264595, 264369, 264685, 264693, 264628, 264563, 284566
2892				UNCLASSIFIED	65274572, 35696286, 29331828, 264110, 265009, 60433438, 265018, 265019, 18108351, 264288, 21906765, 21906766, 21906769, 55811957, 264690, 65274620, 263967, 35695855
				UNCLASSIFIED	264259, 265010, 264682, 18108370, 264555, 264556
		87755985 (5787, 5788) Novel Protein sim. GBank gi 5669015 gb AAD46135.1  - (AF080171) zinc finger protein ZNF232 [Homo sapiens]	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	264259, 265006, 60433438, 52644296, 265011, 264369, 35695917, 18108381, 18108382, 18108388
		86938778 (5789, 5790) Novel Protein sim. GBank gil3924708 emb CAA84646  - (Z35597) Weak similarity with sea squirt nidogen precursor protein (blastp score 71); cDNA EST EMBL:T02069 comes from this gene; cDNA EST EMBL:D76135 comes from this gene; cDNA EST EMBL:D73147 comes from this gene; cDNA EST EMBL:D73147 comes from this gene;	Contains protein domain (PF01437) - Plexin repeat		29331824, 265007, 264762, 264636, 264563
2896		87752122 (5791, 5792) Novel Protein sim. GBank gil4885549 ref NP_005456.1 pPKBG - protein kinase B garmma	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	18108392, 18108394, 18108398, 22278998, 264259, 264259, 29331824, 29146499, 264908, 265007, 265009, 265018, 265019, 264389, 264685, 264689, 21906766, 265021, 264693, 33657182, 264639, 18108384, 18108388, 264567
2897	95413057 (5793, 5794)	95413057 (5793, 5794) Novel Protein sim. GBank gil4502877[ref]NP_001296.1[pCLDN - Clostridium perfringens enterotoxin receptor 1		UNCLASSIFIED	60424179, 56182575, 22278996, 35696286, 22278997, 22278999, 60432049, 264259, 29331824, 29331825, 60424269, 60432289, 29331826, 29331828, 35696052, 264908, 56182435, 265009, 264910, 60170831, 60431735, 60433356, 60433438, 6527444, 55811386, 265018, 18108351, 264448, 264288, 264687, 21906765, 21906766, 21906767, 21906769, 25695763, 264528, 264629, 60431328, 18108374, 55810764, 25811376, 35695423, 3565100764, 26182323, 18108385, 264404, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 2227900
2898	87750340 (5795, 5796)			UNCLASSIFIED	22278995, 264604, 18108385, 264566

2899	180357670 (5797 5798)			INCI ACCIEIED	264764 21006764 264602
200	0423478 (6700 6900)	2000 04233538 (5700 5000) Mana Dratain sim Chark city 604470 cart Charles		ONC CASSIFIED	204104, 21300104, 204032
7007	34233339 (3188, 3600)	Novel Protein sim. Coank gilabota/ulempicAbauts/		glycoprotein	65274572, 56182575, 35696286, 60432049.
	<u> </u>	(118483) SLC/As protein [Homo sapiens]			264259, 29331824, 66714117, 29331826,
					35696052, 29331828, 66712502, 56182435,
					265006, 265007, 265008, 265009, 60433356, }
					264758, 265018, 264764, 264765, 264288,
					264768, 21906764, 21906768, 21906769,
					265020, 264692, 264693, 32833986, 264631,
					83373044, 60432113
2901	_			phosphatase	22278995, 22278997, 22278999, 60432049.
		gij4759272/ref NP_004614.1 pTTC4 - tetratricopeptide			29331822, 29331824, 29331825, 29331827,
_		repeat domain 4		-50	35696052, 33656970, 264910, 265009,
				en c	21906754, 33657084, 87168474, 265010,
					265018, 21906764, 21906765, 21906766,
					21906767, 21906769, 33657023, 264693,
		•			33657109, 33657349, 35696423, 35695855,
					263981, 56182323, 22279002
2902		85745271 (5803, 5804) Novel Protein sim. GBank gil2414615 emblCAB16364  -			264683, 264691
		(299259) hypothetical protein (Schizosaccharomyces			
2903	87606733 (5805 5806)	87606733 (5805-5806) Novel Protein sim GBank nit1079318 high S22241 - XI Ct 2			264887 22278004 264260 20224826
2	(0000 (0000) 00 (000)	protein - African classes from			20224020 264005 52644045 56402425
		Son Donate in the second secon			2930 1020, 204900, 32044040, 30 102433,
					264511, 265017, 265018, 18108351, 264448,
					264683, 264769, 264689, 35695917.
					52644150, 87168518, 60432113, 22279002
2904		86458072 (5807, 5808) Novel Protein sim. GBank		UNCLASSIFIED	52646365, 22278999, 264259, 35696052,
		gij5639823 gb AAD45885.1 AF14367 - (AF143676)			265011, 265017, 264683, 264769, 35695917,
		multispanning nuclear envelope membrane protein nurim			265020, 263967, 18108374, 35695855,
	_	[Homo sapiens]			264637, 264952, 18108385, 18108387
2902		84449926 (5809, 5810) Novel Protein sim. GBank		oncogene	265009, 264681, 264682
		gij728837/spjP39194/ALU7_HUMAN - !!!! ALU SUBFAMILY		ı	
3		SO WARNING ENTRY !!!			
2806	95341051 (5811, 5812)	95341051 (5811, 5812) Novel Protein sim. GBank		UNCLASSIFIED	22278996, 35696286, 22278998, 264259,
		gij4689256jgbjAAD27831.1jAF12185 - (AF121858) sorting	PX domain		60432289, 29331828, 29331830, 66712502,
		nexin 8 [Homo sapiens]			265009, 60170831, 33109954, 264448,
					264683, 264288, 264689, 21906766,
					21906767, 21906768, 55811957, 35695917,
					265022, 52644150, 264691, 33657023,
					264692, 264693, 35695855, 60432113,
					264566
			***************************************		

1807	91211383 (5813, 5814)	2907   91211383 (5813, 5814) Novel Protein sim. GBank gi[1707079 (U80451) - contains strong similarity to a DNAJ-like domain (PS:PS00636) [Caenorhabditis elegans]	Contains protein domain (PF00226) - eph DnaJ domain	фф	52644507, 56182575, 56181686, 22278995, 56994075, 35696286, 60432049, 56182181, 35696052, 64431735, 224595, 55812038, 21906754, 55811386, 2650519, 264682, 264369, 56181562, 21906766, 55811957, 35695917, 265020, 265021, 33657023, 33657109, 60431528, 55811576, 35696423, 35695855, 264638, 22279000
2908	80414246 (5815, 5816)	80414246 (5815, 5816) Novel Protein sim. GBank gil2673917 (AC002561) - putative ATP-dependent RNA helicase [Arabidopsis thaliana]		helicase	265009, 33109954, 18108351, 264766. 265021, 264691, 264692, 18108374, 264556, 264638, 264557, 264558
2909	87420225 (5817, 5818)			eph	264259, 87168474, 265018, 18108365, 264628
2910	86601075 (5819, 5820)	86601075 (5819, 5820) Novel Protein sim. GBank gil4539335 emb[CAB37483.1] - (AL035539) putative protein [Arabidopsis thaliana]			22278995, 264509, 264512, 265007, 33657402, 265017, 264369, 265022, 18108365, 264628
2911	94216615 (5821, 5822)	94216615 (5821, 5822) Novel Protein sim. GBank gil4469187 emb CAB38415.1  - (AL031588) dJ1163J1.3 (novel protein similar to mouse B99) [Homo sapiens]		glucoamylase	52646365, 18108397, 22278995, 22278997, 222789987, 22278998, 22278999, 29331824, 29331825, 52644045, 265006, 265018, 264448, 21906765, 21906766, 21906767, 21906768, 21906769, 55811957, 265021, 18108370, 18108372, 18108372, 18108374, 22279000
2912	87731803 (5823, 5824)	2912   87731803 (5823, 5824) Novel Protein sim. GBank gil4929637[gb]AAD34079.1 AF15184 - (AF151842) CGI-84 Involucrin repeat protein [Homo sapiens]	Contains protein domain (PF00904) - Involucrin repeat		52645156, 264092, 60432049, 264259, 52645080, 29331824, 29331825, 66712502, 33109954, 264760, 264683, 264288, 264686, 265021, 264693, 18108368, 263976, 264404
2913		87713823 (5825, 5826) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]	Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	UNCLASSIFIED	52644507, 52645156, 52646842, 56182575, 35696286, 22278997, 264259, 52645080. 29331827, 3569628. 22331828, 264828. 52644045, 56182435, 55812038, 52646317, 21906754, 2564296, 87168474, 265017, 265018, 265049, 18108351, 264682, 264686, 21906768, 21906762, 21906768, 25694150, 27486261, 27486262, 27486265, 35695855, 52644332, 22279000, 22279002, 264563
2914	87797300 (5827, 5828)				264557

2015	88081072 /5830 5830)	2015 IRROR1072 (FRESO FRESO) Novice Provision sim Charles	ten (050000) diamete dictore societare	400	99590010 951190 560190 99190 993190	Γ
	(2000 10100)	aliS174485trefine monator	Lection Catone domain		50546842 21906767 21908768 56180575	<del>.</del>
		(macrophage mannose receptor family)			22148629 35696917 22278996 22278997	
					22278998, 265021, 22278999, 52644150.	
		-			264691, 264259, 60432049, 264692,	
					52645129, 33657109, 33657182, 29331827,	
					27486261, 35696052, 29331828, 27486262,	
					27486264, 27486265, 33657349, 29146498,	
					29146499, 264906, 264907, 18108370,	
					264908, 18108372, 52644045, 18108374,	
					56182435, 35695855, 264112, 264510,	
					265008, 60432229, 264593, 60433356,	_
					56182323, 18108382, 55812038, 18108385,	_
					33109954, 21906754, 33657084, 87168518,	
					265010, 265011, 60432113, 265017, 265018,	<u></u>
					22279000, 265019, 55811150, 264681.	
					18108351, 264763, 264448, 264683, 264566,	,. <u>.</u>
0,00	_	17 10700 4 4 Cl. 18 17 10 C C C C C C C C C C C C C C C C C C			18108354, 264369, 264288, 264766	Т
2816	_	9533779U (5831, 5832) Novel Protein Sim. GBank gij5104851 dbjjBAA80165.1		dehydrogenase	52645156, 65274572, 22278994, 22278995,	_
		(AP000061) 305aa long hypothetical dTDP-4-			35696286, 22278996, 22278997, 22278998,	
		denydromamnose reductase (Aeropyrum pernix)			22278999, 264259, 29331822, 29331824,	
					29331825, 66714117, 60432289, 29331826,	
					29331827, 29331828, 33656970, 264509,	
					264906, 29331830, 52644045, 264909,	_
					36182433, 60170831, 264392, 264393,	
					3365/402, 60433356, 52646317, Z1906/54,	
					33109934, 3383/004, 32044296, 83838342,	
					265011, 265017, 265018, 265019, 18108351 264448 264288 52644230 21006265	<u>:</u>
					204449, 604200, 32044223, 21300103,	_
					Z1906/66, Z1906/67, Z1906/68, Z1906/69,	
					33011337, 33033317, 203021, 203022, 62644460, 32663033, 32663400, 32663403	
			-		32044(30, 3303/023, 3303/109, 3303/102, 37486364 37486362 37486364 36606763	
					27400201, 27400202, 27400204, 33033703, 18108376, 55811576, 35696423, 65274791	
					35695855, 52644332, 264557, 264638,	
					56182323, 18108387, 87168518, 22279002,	
					264482	
2917	87454546 (5833, 5834)	87454546 (5833, 5834) Novel Protein sim. GBank gil3169065 emb CAA19260.1 -		UNCLASSIFIED	60433438, 264602, 264682, 87168518,	Г
		(AL023704) putative translocation elongation factor-Tu fa			60432113	
2018	85600520 (5835 5836)	RESOURCE (SAME EDGE) Name Destrict Chart allegations			26.4630	Т
9	(2022, 2020)	inoval rioteti sim. Obalit gijosez topnijosooso - prypothetical protein YKL201c - yeast (Saccharomyces rervisiae)			000007	
2919	87641497 (5837, 5838)	87641497 (5837, 5838) Novel Protein sim. GBank gi[2564955 (AF030001) -			66714117, 66712502, 263981	П
		unknown [Mus musculus]				$\Box$

2920	2920 87769523 (5839, 5840)	[(			35696286, 22278997, 264259, 52645080.
					29331824, 29331826, 29331827, 264828,
-					264909, 56182435, 264511, 264758,
					33109954, 21906754, 52644296, 265010,
					265011, 264601, 265017, 265019, 264681,
					264687, 21906767, 265021, 52644150,
					264690, 264691, 264692, 264693, 33657109,
					33657182, 27486262, 27486264, 27486265,
_					35696423, 35695855, 264632, 264636,
					264637, 264638, 56182323, 60170394,
_					18108385, 87168518, 60432113
2921 9	31639982 (5841, 5842 <u>)</u>		Contains protein domain (PF00787) -		35696286, 22278997, 264091, 264092,
		gi[4580013]gb AAD24202.1 U83194 (U83194) TRAF4-	PX domain		264094, 264259, 29331822, 29331824,
		associated factor 2 [Homo sapiens]			29331826, 29331827, 35696052, 29146498,
					264104, 264105, 264107, 264509, 264110,
					264112, 264512, 60433356, 21906754,
					87168474, 265017, 18108351, 264288.
					21906765, 21906766, 21906767, 21906769.
					35695917, 265021, 263974, 18108374,
					263976, 263977, 18108376, 264555, 263981,
_					56526486, 87168518, 22279000, 22279002
2922 8	17749762 (5843, 5844)	87749762 (5843, 5844) Novel Protein sim. GBank gil4589514 dbj BAA76779.1  -	Contains protein domain (PF01074) - kinase	kinase	264906, 264909, 264511, 265006, 265008.
		(AB023152) KIAA0935 protein [Homo sapiens]	Glycosyl hydrolases family 38		264593, 33657402, 60174639, 18108351
					264763, 21906765, 29148627, 35695917,
					264692 264629 261978 45811576
					204092, 204023, 2033/0, 300113/0, 35695855 264555 264558 46182323
$\overline{}$					60170394, 22279000, 264486
2923 9	5337799 (5845, 5846)	95337799 (5845, 5846) Novel Protein sim. GBank ail4835268lemblCAB42898 21 -	Contains protein domain (PE00169) - struct	strict	26448R 1810R397 22278995 22278995
_	<u>, , , , , , , , , , , , , , , , , , , </u>	1783844) 4137E16 4 (cimilar to mouro p1460io proteio)	Columnia Proventi Collingini (r. 1 00 103) - 2	20116	204400, 10100397, 22270339, 22270390,
		(Loso44) 0337 E10.4 (Similar to mouse pritorip protein)	TH domain	•	22278997, 22278998, 22278999, 29331825,
_					29331826, 29331827, 29331830, 264511,
					265009, 33657402, 265011, 265017, 265018,
					264683, 18108354, 21906765, 21906767,
					21906768, 21906769, 52644150, 264691,
					264692, 33657109, 263974, 18108376,
					264631, 264636, 18108385, 18108387,
づ					22279000, 264563, 264566
2924 8	17791967 (5847, 5848)	87791967 (5847, 5848) Novel Protein sim. GBank gil2133095[pir] S72254 -	Contains protein domain (PF00444) - ribosomalprot	ribosomalprot	265017, 264628, 20281152, 264556
		ribosomal protein L36, mitochondrial - yeast	Ribosomal protein L36		
3000	2000400 40000	(Saccharomyces cerevisiae)			
	5090120 (5849, 5850)	95090120 (5849, 5850) Novel Protein sim. GBank gi[2388986 emb CAB11718  -		UNCLASSIFIED	56182575, 35696286, 264259, 60432289,
		(Z98980) actin associated protein [Schizosaccharomyces			29331827, 264508, 52644045, 264910,
		[bombe]			264591, 60432229, 55812038, 21906754,
					264681, 264448, 264683, 264288, 264685.
					52644229, 264689, 21906765, 21906766,
					21906768, 21906769, 265021, 265022,
					60170615, 264692, 33657023, 264693,
					33657109, 35696423, 65274791, 56182323

2928	95343003 (5851 5852)				
2827	7	80408018 (5853, 5854) Novel Protein sim, GBank oil 283032 Inicil S22456.			29331628, 2650111, 264/68, 264689
	_	hydroxyproline-rich glycoprotein - perennial teosinte			204/04, 204266, 204630, 20463/
2828				UNCLASSIFIED	264559
2929		91622920 (5857, 5858) Novel Protein sim. GBank gij3413320 embjCAA06915		UNCLASSIFIED	264569, 264489, 22278994, 35696286,
		(AJ006215) CMP-N-acetyineuraminic acid synthetase [Musmusculus]			22278996, 22278998, 22278999, 264094, 264259, 52645080, 29331822, 29331824, 66714117, 29331825, 29331826, 29331827, 35696052, 33656970, 264109, 29331830,
					52644045, 265009, 33109954, 52644296, 87168559, 264760, 264762, 264448, 264764,
					.264288, 264766, 264768, 21908765, 21906766, 21906768, 21906769, 35695917,
					264691, 33657023, 264693, 33657109, 18108374, 263976, 35696423, 35695855,
					263981, 22279000, 22279002, 264567, 264486
2930	95302755 (5859, 5860)			UNCLASSIFIED	56182575, 56181686, 35696286, 22278996,
					ZZZ/8996, ZZZ/8999, Z64Z59, Z93318Z5, ( 60432389, 20331828, 264905, 52644045
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					60432229, 60433356, 87168474, 265010,
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_					264448, 264683, 264288, 264766, 21906765,
					21906769, 35695917, 60170615, 33657023,
					33637109, 264626, 18108370, 18108372, 35696423, 35695855, 264556, 56182323
	$\overline{}$				60432113, 264567
2931		3786433 (AF098505) - similar	Contains protein domain (PF00471) - UNCLASSIFIED	UNCLASSIFIED	52645156, 22278997, 22278998, 29331822,
		to Arabidopsis thaliana male sterility protein 2 (SW.Q08891) Ribosomal protein L33	Ribosomal protein L33		52645080, 29331824, 60432289, 33656970,
		[Caenomabditis elegans]			60433356, 60433438, 33109954, 21906765.
					21906766, 21906767, 21906768, 265020,
					0204410U, 3300/UZ3, 3300/109, 3300/16Z,
	_	-			27.400200, 33089423, 33083303, 204333, 87168518, 60432113, 264566
2932	_				264906, 264907
2933		91720776 (5865, 5866) Novel Protein sim. GBank gij3378056 (AF017777) - helicase		helicase	264488, 18108392, 56182575, 22278999,
		Lorosopniia meianogasteri			264091, 264259, 29331825, 60432289,
_					29331627, 264508, 52644045, 56182435,
_					60433438 3400874 366047 364683
_				•	264288, 52644229, 21908765, 21908766.
					21906768, 21906769, 265022, 52644150.
					33657023, 33657109, 27486265, 264635,
					264636, 60170394, 56182323, 18108385,
					60432113, 264565, 264566, 264567

2934	2934 86576025 (5867, 5868)				22278997, 22278999, 29331824, 33657402
					264691, 27486262, 264628, 87168518.
					22279000
2835	86410579 (5869, 5870)			UNCLASSIFIED	56182575, 22278995, 60433356, 33657402,
					264758, 33109954, 21906754, 265018,
					265019, 264448, 264769, 21906764,
					21906765, 265021, 264692, 33657023,
					33657109, 33657349, 55810764, 22279000
2936		87605863 (5871, 5872) Novel Protein sim. GBank gil4153862 (AC005065) -	Contains protein domain (PF00856) - nuclease	nuclease	22278997, 29331827, 29331828, 265009,
		determined by GENSCAN prediction and spliced EST;	SET domain		265017, 264605, 265020, 55811576,
	Т	match to EST R84329 (NID:942735) [Homo sapiens]			18108387, 60432113, 264563
2937		94853096 (5873, 5874) Novel Protein sim. GBank		UNCLASSIFIED	56994075, 22278999, 264259, 60432049,
		gi[5174409 ref NP_006101.1 pCD2B - CD2 antigen			29331822, 56182181, 29331827, 29331828,
		(cytoplasmic tail)-binding protein 2			264906, 264908, 264909, 56182435, 265006,
					264512, 264910, 60170831, 60433356,
					265011, 265018, 18108351, 264448, 264288,
					264766, 52644229, 21906765, 29148784,
					65274791, 264556, 56182323, 60170394,
	$\overline{}$				264558, 60432113, 264565, 264486, 264567
2938		95419773 (5875, 5876) Novel Protein sim. GBank gij3319990jembjCAA76720j -	Contains protein domain (PF00179) - ubiquitin	ubiquitin	264488, 56182575, 22278996, 35696286,
		(Y17267) ubiquitin-conjugating enzyme [Mus musculus]	Ubiquitin-conjugating enzyme		22278997, 22278998, 22278999, 264490,
					264259, 29331822, 29331824, 66714117,
					29331827, 35696052, 264107, 264905,
					66712502, 52644045, 56182435, 264511,
					265008 265009 60432229 33657402
					60433438, 55812038, 21906754, 85658542
					265010 265011 B7168550 265017 265049
_					255010, 253011, 8/158339, 263017, 263018, 1
					202013, 204001, 204200, 204003, 21300703,
					21906/67, 21906/68, 55811957, 35695917,
					265020, 60170615, 264690, 264691, 264692,
					33657023, 264693, 65274620, 33657109,
					18108370, 18108374, 263976, 35696423,
					35695855, 264555, 264556, 18108381,
					56182323, 60170394, 83373044, 18108385,
					56526486, 60432113, 22279002
2939	87786622 (5877, 5878)	2939   87786622 (5877, 5878)   Novel Protein sim. GBank gij3979900 emb CAA99909  -	Contains protein domain (PF00400) -	ATPase_associated	Contains protein domain (PF00400) - ATPase_associated 264907, 265018, 264681, 264685, 264686
		(275547) similar to WD domain, G-beta repeat; cDNA EST	WD domain, G-beta repeat		
		yk3/10/.5 comes from this gene; cDNA EST yk312h1.5			
		comes from this gene; cDNA EST yk465d5.5 comes from			
		this gene; cDNA EST yk472c4.5 comes from this gene;		-	
		CUNA ESI YKZYZIB			

8	2940   95011103 (5879, 5880)		1	UNCLASSIFIED	22278996, 29331822, 29331824, 66714117,
					29331826, 29331828, 264905, 264908,
					66712502, 29331830, 265011, 265017.
					264764, 264369, 21906766, 21906767,
					33657023, 33657109, 32833986, 18108374,
					18108377, 264634, 83373044, 18108385,
					18108387, 264566
2941	21423370 (5881, 5882)	21423370 (5681, 5682)  Novel Protein sim. GBank gi 3413872 dbj BAA32300  -  (AB007924) KIAA0455 protein [Homo sapiens]		UNCLASSIFIED	264557
2942	87430203 (5883, 5884)	87430203 (5883, 5884) Novel Protein sim. GBank		głycoprotein	264910, 265010, 264768
		gi 1172845 sp P46629 RB25_RABIT - RAS-RELATED		•	
2943		95314504 (5885, 5886) Novel Protein sim. GBank	)	collagen	60432049, 264259, 60432289, 29331827,
		gil4929653 gb AAD34087.1 AF15185 - (AF151850) CGI-92		<del></del>	29146498, 265008, 264593, 60433356,
		protein [Homo sapiens]			60433438, 265010, 265011, 265017, 265018,
					264683, 264766, 18108381, 65274727.
					60432113, 264567
2944			Contains protein domain (PF00415) - /	ATPase_associated	Contains protein domain (PF00415) - ATPase_associated   56994075, 22278998, 60432049, 264259,
		(ALCASOOU) 1-acylcetor-3-priospriate acylitatisterase-like	regulator of chomosome		23331022, 23331024, 00424203, 00434203, 20331826, 20331828, 264005, 264007
		_	College Salion (1900)		23031010, 23031010, 204300, 104300; 52644045, 264909, 264511, 265006, 265009
		-			264594 21906754 R7168559 264603
					265017 265018 18108351 264682 264766
	•				264687 264689 21906765 21906766
					21906767 21906768 21906769 265021
					61300101, 21300100, 21300103, 203021, 60170615, 52644150, 264690, 264691
					30110013, 32044130, 204030, 204031.
					33657023, 264692, 264693, 33657109.
					3365/182, 3365/349, 181083/0, 181083/4,
					18108377, 55811576, 35696423, 35695855,
					264635, 264555, 264556, 56182323,
					60170394, 264558, 264559, 83373044,
					56526486, 87168518, 60432113, 22279002,
					264482, 264563, 264484, 264567
2945			Contains protein domain (PF00096) - UNCLASSIFIED		60424179, 22278995, 22278996, 22278998.
		gif728631[sp[P39188]ALU1_HUMAN - !!!! ALU SUBFAMILY  Zinc finger, C2H2 type	Zinc finger, C2H2 type		22278999, 264259, 56182181, 29331824,
		J WARNING ENTRY !!!!			60424269, 60432289, 35696052, 264908,
					265006, 60433356, 55812038, 264759,
	_				55811386, 265018, 264681, 18108351,
					264448, 264683, 264369, 264288, 264687.
					30101304, Z1800101, Z1800100, Z1800103,
					SOBSOBIL, ZGOUZU, ZGOUZI, ZG4G93. SO431528 KE810764 RESOGA23 RESOGRES
					00451360, 03010704, 03030423, 03033003,
					20405U, 00170584, 05375044, 22279000,
1					204300, 204307

2046	194317315 (5891 5802)	2046   04317315 (5801 5802)   Navel Destein ein Chank		INC. ACCICION	DEALOR DEADED DEAEND DEALEND
2	(1000)	ALEGATORNICAL SUM COMME TARGETORS (ACCORDED)			264007 264000 264640 266003
		31944 1934 1994 1995 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			7005011, 204503, 204310, 204311, 203007,
_		peroxisomal membrane protein PMP 24 [Homo sapiens]			[284512, 264910, 264591, 264593, 18108351,
					264764, 264288, 264684, 264769, 265021,
					264692, 33657109, 264628, 264629,
					18108374, 264631, 264634, 264636, 264637,
_					18108380, 264638, 264639, 83373044
					264565, 264566, 264486, 264567
2947	87362952 (5893, 5894)	87362952 (5893, 5894) Novel Protein sim. GBank gij3540281 gb AAC34383.1  -		UNCLASSIFIED	22278995, 22278996, 22278997, 22278999,
		(AF056116) All-1 related protein [Fugu rubripes]			29146498, 264508, 29331830, 265007,
					265008, 265009, 60432229, 21906754,
					265010, 265017, 265019, 264766, 264685,
					21906765, 21906766, 21906767, 21906768.
					21906769, 265020, 264628, 18108370,
					264629, 264630, 18108387, 60432113
2948	87626527 (5895, 5896)	87626527 (5895, 5896) Novel Protein sim. GBank gi 5566614 gb AAB65654.2  -			52646842, 22278995, 264259, 29331824,
		(AF001533) mitogen-induced [Mus musculus]			29331825, 29331827, 29331830, 264909,
					265007, 265009, 265019, 264763, 264684.
					264288, 264685, 264686, 21906767, 264691,
			•		264692, 264693, 18108374, 55811576,
					18108385, 22279002, 264563, 264567
2949		88175545 (5897, 5898) Novel Protein sim. GBank gi 2132923 pir  S67133 - probable		UNCLASSIFIED	22278996, 22278997, 60432289, 29331826,
		membrane protein YOR240w - yeast (Saccharomyces			29331827, 29331828, 35696052, 29146499,
		cerevisiae)			264104, 264107, 264905, 66712502, 264908.
					60433356, 60433438, 87168559, 264764,
					52644229, 56181562, 21906767, 21906768.
					21906769, 265022, 60170615, 33657023,
					35696423, 263981, 264558, 60432113.
	·				22279002
2820	95086870 (5899, 5900)	2850   95086870 (5899, 5900)   Novel Protein sim. GBank	Contains protein domain (PF00883) - peptidase	peptidase	264488, 35696286, 264259, 35696052,
		gilabo iuzispir 34629 TUJb_CAEEL - PUTATIVE	Cytosol aminopeptidase ramity		264907, 263007, 264910, 263017, 263018,
		AMINOPEP IIDASE 2K353.6 IN CHROMOSOME III			264288, 264768, 35695917, 265020,
					18108362, 18108370, 18108379, 35696423,
-	-,				65274791, 35695855, 264556, 56526486.
					264486
2951	87392357 (5901, 5902)	87392357 (5901, 5902) Novet Protein sim. GBank gij4688902 emb CAB41450.1  -  (AJ238248) centaurin beta2 tHomo saniens			264693
ĺ	ţ				

2952	95329952 (5903, 5904)	2952   95329952 (5903, 5904) Novel Protein sim. GBank gil5596693 emb CAB51405.1  - (AL096881) hypothetical protein [Homo sapiens]	Contains protein domain (PF00650) - transcriptfactor	transcriptfactor	264687, 52645156, 21906766, 21906769, 22278996, 265020, 264690, 60432049, 264259, 264690, 60432049, 264259, 264690, 60432049, 29331825, 264692, 29331827, 35696052, 2748626, 264508, 264905, 20331830, 264908, 264907, 29331830, 264908, 264900, 264635, 264511, 265008, 264909, 35695855, 264636, 60432229, 264638, 60433356, 264638, 60432229, 264760, 264563, 26279000, 22279000, 26279002, 264488, 264288
2853		88093575 (5905, 5906) Novel Protein sim. GBank gil119522 sp P10658 SERC_RABIT - PROBABLE PHOSPHOSERINE AMINOTRANSFERASE (PSAT) (ENDOMETRIAL PROGESTERONE-INDUCED PROTEIN) (EPIP)	Contains protein domain (PF00266) - UNCLASSIFIED Aminotransferases class-V	UNCLASSIFIED	18108398, 56994075, 22278996, 29331822, 29331824, 29331825, 29331827, 35696052, 264508, 264905, 264907, 264510, 264591, 264591, 264592, 264907, 264510, 264758, 5264634, 21906754, 264750, 264681, 18108351, 264764, 264369, 264288, 264681, 18108351, 264764, 264369, 269288, 21906768, 21906766, 21906766, 21906767, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 3665917, 33657023, 18108374, 263978, 35695855, 264637, 264638, 87168518, 264482, 264563, 264565
2954		88086288 (5907, 5908) Novel Protein sim. GBank gil4885261 ref NP_005251.1 pGDF9 - growth differentiation factor 9	Contains protein domain (PF00019) - Transforming growth factor beta like domain	lgí	29331822
2955		87698426 (5909, 5910) Novel Protein sim. GBank gij3452473 (AF084205) - serine/threonine protein kinase TAO1 [Rattus norvegicus]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264259, 29331822, 29331824, 29331825, 29331825, 29331826, 35696052, 264908, 52644045, 264512, 60432229, 265018, 265019, 55811150, 264769, 21906767, 21906768, 21906769, 21906769, 265021, 60170615, 55810764, 264567
2956		85789745 (5911, 5912) Novel Protein sim. GBank gil4689254[gb]AAD27830.1 AF12185 - (AF121857) sorting   nexin 7 [Homo sapiens]	Contains protein domain (PF00787) - PX domain		22278996, 264259, 29331827, 264908, 21906768
2957	80933301 (5913, 5914)	90933301 (5913, 5914) Novel Protein sim. GBank gi 4503023 ref NP_000089.1 pCPT2 - carnitine palmitoyttransferase II precursor		cadheiin	22276999, 264259, 29331824, 29331827, 265008, 264595, 264758, 265010, 265011, 264448, 264763, 264683, 264288, 264685, 18108357, 29148629, 264690, 18108362, 264699, 18108362, 264693, 18108370, 60431528, 18108374, 264634, 18108381, 56182323, 18108382, 18108382, 264482, 264487
2958		87440014 (5915, 5916) Novel Protein sim. GBank gil4240257[dbj BAA74907.1  - (AB020691) KIAA0884 protein [Homo sapiens]			264595, 264596, 264681, 264369, 264629, 264631, 264567

2959   95109420 (5917, 5918) Novel Protein sim. GBank gij988221 (U33005) - Tbc1 [Mus   Contains protein domain (PF00566) - oncogene musculus]
95413416 (5921, 5922) Novel Prolein sim. GBank gij5596646 emb CAB05177.2  - Contains protein domain (P (Z82266) predicted using Genefinder; similar to WD domain, WD domain, G-beta repeat G-beta repeats [Caenorhabditis elegans]
95313464 (5925, 5926) Novel Protein sim. GBank gil4240223 dbj BAA74890.1  - (AB020674) KIAA0867 protein [Homo sapiens]

2965	80384762 (5929, 5930)	2965  80384762 (5929, 5930) Novel Protein sim. GBank		transprintfactor	201260 202100 201200 201200
		gil4885447[ref]NP_005452.1[pKRML - Kreisler (mouse) maf- related leucine zioner homologi			264907, 264908, 264909, 264511, 265008.
		Room and a second secon			264910, 264591, 264593, 264594, 33657402, 265011, 264760, 264762, 264764, 264288
				_	264685, 264766, 264692, 33657109, 264628,
					264629, 35695855, 264630, 264631, 264632,
					264634, 264635, 264636, 264637, 264638,
2000	04775740 (E024 E022)				264639, 264563, 264567, 18108391
7 200	91123240 (3331, 3332)	9 (123249 (3331, 3332) Novel Pfotein sim. GBank gij5262751jembjCAB45690.1[-			60432289, 264682, 264448
		(Xenopus laevis)			
2967	94658303 (5933, 5934)	Novel Protein sim. GBank gil624225 (U19181) - Rabin3		LINCI ASSIFIED	2644RR 26450R 254500 26400R 264000
		[Rattus norvegicus]			264511, 264910, 264594, 264758, 85658542.
					264762, 264764, 265021, 264556, 18108381,
2000	2003, 255000				264564, 264486
7.000	(0580, 0830, 0830)		Contains protein domain (PF00097) -		264687, 52645156, 21906765, 52646365,
		18.1[AF15188 - (AF151881) CGI-123	Zinc finger, C3HC4 type (RING		21906767, 18108398, 35696423, 22278996,
		protein [Homo sapiens]	(inger)		35696286, 22278997, 265020, 22278999,
					265021, 265022, 264093, 264636, 264690,
					52644150, 264259, 33657023, 52645080,
					264693, 29331822, 56182181, 29331824,
					66714117, 29331825, 33109954, 52645129.
					29331826, 21906754, 33657182, 29331827.
					29331828, 35696052, 27486262, 87168518,
		-			87168474, 265010, 87168559, 265018.
					22279000, 265019, 22279002, 264563,
					18108351, 264906, 264907, 264448,
6962	95310957 (5937 5938)	2969 195310957 (5937 5938) Novel Protein sim GRank			66712502, 264566, 264369, 264288
		ū		uda	52646842, 22278996, 22278998, 22278999,
		SISSENTASISPIOZANOAPINOSA, OULON - I HERMOSOME,			60432049, 264259, 29331824, 29331825,
		ALTER SUBGINI (CHAPERONIN ALTER SUBGINI)			29331826, 29331828, 264509, 264909,
					52644045, 56182435, 265009, 60433438,
					55812038, 21906754, 265011, 87168559,
					265018, 265019, 264448, 264288, 264369,
					52644229, 21906766, 21906768, 21906769,
-					29148784, 265020, 265021, 52644150,
					264691, 33657109, 18108374, 56182323,
2970	88088071 (5939, 5940)	88088071 (5939, 5940) Novel Protein sim. GBank gil3165407 (AC004755) .	Contains protein domain (PE00046) - homeobox		00170394, 87108318, 60432113, 22278000
			Homeobox domain	Youngonia	

264488, 56182575, 35696286, 56994075. 29331824, 29331826, 29146499, 264508. 264905, 264907, 264112, 264910, 21906754. 87168559, 265018, 265019, 18108351, 264689, 21906765, 21906767, 21906768, 265020, 265021, 60170615, 18108364, 264628, 264629, 18108374, 264636, 264556, 264558, 83373044, 18108384, 18108385,	265017, 35695917, 265021, 33657109, 22279002, 264563	29331822, 264692, 33657349, 55811576, 264563	18108392, 52644507, 56182575, 56181686, 22278993, 22278996, 35696286, 22278997, 22278998, 2569286, 22278997, 22278998, 264259, 52645080, 29331824, 29331824, 29331824, 29331824, 2565020, 2644045, 265007, 264910, 265009, 60433438, 33109954, 21906754, 55811386, 52644296, 87168474, 87168559, 265017, 265018, 265019, 18108351, 264448, 264369, 264288, 52644229, 18108359, 21906765, 21906767, 21906768, 35695917, 265020, 265021, 52644150, 264691, 264692, 33657023, 27486262, 27486264, 35695763, 18108370, 18108376, 55810764, 58811576, 35696423, 35695855, 264537, 5264432, 264581, 22278000, 264482, 264487	29331824, 29331827, 29331828, 264910, 85658542, 265011, 265018, 264448, 264288, 264769, 21906767, 265020, 264691, 264559, 83373044	22278996, 264906, 265007, 265010, 265011, 265017, 265017, 264689, 18108351, 264685, 264689, 18108370, 264639, 18108385	264509, 264288
tm <b>7</b>	kinase	UNCLASSIFIED	UNCLASSIFIED	ubiquitin	transport	UNCLASSIFIED
				Contains protein domain (PF00632) - ubiquitin HECT-domain (ubiquitin- transferase).		
94196930 (5941, 5942) Novel Protein sim. GBank gil728837{sp P39194 ALU7_HUMAN - IIII ALU SUBFAMILY SQ WARNING ENTRY IIII	86625943 (5943, 5944) Novei Protein sim. GBank gi 728836 sp P39193 ALU6_HUMAN - III! ALU SUBFAMILY SP WARNING ENTRY IIII		91673002 (5947, 5948) Novel Protein sim. GBank gil786117 (L41834) - nuclear protein [Ensis minor]	95325213 (5949, 5950) Novel Protein sim. GBank gij3880812 emb CAA19508  - Contains protein domain (AL023839) similar to HECT-domain (ubiquitin-transferase).; HECT-domain (ubiquitin-cDNA EST yk480d10.5 comes from this gene	87771202 (5951, 5952) Novel Protein sim. GBank gi 5679136 gb AAD46874.1 AF16093 - (AF160934) BCDNA.LD14189 [Drosophila melanogaster]	91725254 (5953, 5954) Novel Protein sim. GBank gil5262751 lemb CAB45690.1  -  (AJ243177) Xenopus RPA interacting protein alpha  [Xenopus laevis]
1 94196930 (5841, 5942)						
2971	2972	2973	2974	2975	2976	2977

2978	87332059 (5955, 5956)	2978   87332059 (5955, 5956) Novel Protein sim. GBank gil746549 (U23522) - No	Contains protein domain (PF00480) - UNCLASSIFIED	UNCLASSIFIED	22278995, 22278996, 22278997, 22278999.
			<b>6</b>		56182435, 265006, 265007, 265009,
					60433356, 60433438, 21906754, 265010,
					265011, 265017, 265018, 265019, 264288.
					264685, 264688, 21906765, 21906766,
					21906767, 21906768, 21906769, 265020,
					265021, 265022, 35696423, 264639,
					60432113, 22279000, 22279002
2979		91725256 (5957, 5958) Novel Protein sim. GBank gi 5262751 emb CAB45690.1  -		complement	264488, 65274572, 56994075, 22278999.
		(AJ243177) Xenopus RPA interacting protein alpha			264093, 29331822, 29331824, 264288,
		[Xenopus laevis]		K	55811957, 33657023, 33657109, 18108370,
	_				55811576, 56182323, 60432113, 264482
2980	86296600 (5959			***	265009, 21906767, 263981, 22279000
2981				UNCLASSIFIED	264629, 264564
2982		95303675 (5963, 5964) Novel Protein sim. GBank			22278995, 56994075, 22278996, 22278997,
		gi 4929767 gb AAD34144.1 AF15190 - (AF151907) CGI-149			22278998, 22278999, 264092, 29331824,
		protein [Homo sapiens]			29331827, 29331828, 264905, 264591,
	٠				264592, 264594, 264595, 264596, 33657084,
					264448, 21906765, 21906766, 21906767,
				•	21906768, 21906769, 265020, 265022,
					18108365, 33657182, 33657349, 35696423,
					83373044, 22279000, 22279002
2983		91725258 (5965, 5966) Novel Protein sim. GBank gij5262751 emb CAB45690.11			60424179, 52646842, 18108398, 22278997,
	-	(AJ243177) Xenopus RPA interacting protein alpha			264093, 60432049, 264259, 29331822,
		[Xenopus laevis]			60432289, 33656970, 264905, 52644045,
					265006, 60431735, 87168474, 265018,
		-			265019, 18108351, 264448, 21906765,
					21906768, 35695917, 33657023, 52645129,
					18108370, 35696423, 83373044, 56526486,
					60432113, 264404, 22279002
29 <u>8</u> 4	94136467 (5967, 5968)	2984   94136467 (5967, 5968) Novel Protein sim. GBank gil2393734 (AC002542) - similar		ATPase_associated	
		to C. elegans F11A10.5; 80% similarily to Z68297 [PfD:o1130619) [Homo sapiens]			
2985	87099072 (5969 5970)	87099072 (5969 5970) Novel Protein sim GRank dil10316010iri1522128 . finas		LINCI ACCIEIED	254010 55812018 56181562 55811057
2	(0.00', 0.00', 0.00')	prover notein sim. Geain gij 103 100 pii j322 120 - miger Profein unkernet fauit (in (Oceanhile metacocarter)		CACCASSIFIED	204310, 33812030, 38181302, 33811337, 384838 85810764 384833 384838
					60432113
5986	86284861 (5971, 5972)				55811957, 264566
2987				UNCI ASSIETED	264369
	7 ' '			טואניהטטיוט	204303

129BE	8  95357753 (5975 5976	Novel Drotoin aim Charle allegacon and allegacon			
	0.00 CO	(AF077207) Howel Protein sim, GBank gil4679028 gb AAD27002.1  - (AF077207) HSPC021 [Homo sapiens]		UNCLASSIFIED	264488, 62274572, 22278995, 22278996, 22278997, 22278999, 264092, 264094, 264259, 264092, 264094, 264259, 264092, 264094, 264269, 264328, 29331824, 29331826, 60432289, 35696052, 29331828, 264107, 264905, 264908, 66712502, 264828, 264909, 56182435, 265006, 265007, 265008, 60170831, 60432229, 264593, 60433356, 265011, 87168559, 265017, 265018, 264682, 265011, 87168559, 265017, 265018, 264429, 265011, 265021, 265022, 26244150, 265021, 265022, 26244150, 265021, 265022, 26244150, 264680, 33657023, 6527450, 265062, 265022, 265021, 265022, 263697, 265021, 265022, 263697, 265021, 265022, 265021, 265022, 265021, 265022, 265021, 265022, 265021, 265022, 265021, 265022, 265021, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265022, 265
					264636, 264637, 263981, 264638, 56182323, 83373044, 6043213, 22279000, 264563, 834564, 264665, 264665
6067		91423118 (3977, 3978) Novel Protein sim. GBank gi 113671 sp P23964 ALUF_HUMAN - !!!! ALU CLASS F WARNING ENTRY !!!!		kinase	22278996, 22278997, 264905, 284511, 60170831, 264593, 265019, 21906765,
2880			Contains protein domain (PF00071) - oncogene Ras family	oncogene	21906767, 21906768, 18108374 265007, 264512, 18108351, 264288, 264689, 265020, 264691, 33657023, 33657109
2991 2991	84325361 (5981, 5982) 85425164 (5083, 5084)				264563
283 283					264259, 265019, 264689, 18108385
9				UNCLASSIFIED	264488, 29331822, 265017, 264761, 21906769, 65274701, 263081, 264665
4887		94136634 (5987, 5988) Novel Protein sim. GBank 91 2496549 sp Q50658 YU02_MYCTU - HYPOTHETICAL 29.7 KD PROTEIN CY339.02		transport	22278994, 22278995, 56994075, 22278997, 22278999, 264259, 29331822, 29331828, 29331828,
					265006, 265009, 264910, 33109954, 87168474, 87168559, 265018, 265019, 264448, 264288, 21906766, 21906767, 21906768, 21906769, 265021, 265022, 33657023, 264693, 35695855, 83373044,
2995		87591070 (5989, 5990) Novel Protein sim. GBank gi[2734081 (AF000195) - similar to oxysterol-binding proteins [Caenorhabditis elegans]			18108385, 22279000, 264565, 264566 264905, 264907, 265019, 18108351, 264683
0 6 6 7		91013798 (5991, 5992) Novel Protein sim. GBank gi[2829912 (AC002291) - Similar ATP-dependent RNA Helicase [Arabidopsis thaliana]	Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase		65274572, 35696286, 264259, 29331824, 35696052, 29146499, 264508, 264907, 265007, 265008, 60433438, 18108348, 265017, 264681, 264683, 264288, 264766, 264769, 264689, 35695917, 60170615, 366573, 26428
					18108381, 18108382, 18108388, 264484

1000	11 002 00021 011 50050			
	01041110 (3883, 3884)	(1992), 1999, 1999, 1999, Mover Protein Smr. GBank gli4589552[ab] BAA75848.1  - (1995), 1999, May 1999, Ma	homeobox	264488, 56182575, 264259, 66714117, 29331826, 35696052, 264508, 264509,
				264907, 264908, 265006, 87168474, 265019.
				264448, 264682, 264685, 264766, 21906764,
				21906766, 21906768, 21906769, 27486261,
				18108374, 35696423, 264634, 264635,
2000	10002 2002 10000			264636, 264557, 18108385, 87168518
0667	(0880, 0880) 10008000	ocusoso I (osaso, osaso) More Frolein sim. Csank gilj3947889[emb](AAA22252]	UNCLASSIFIED	52646365, 22278997, 264508, 264906,
		(ALUS4304) CUNA EST ykt3509.3 comes from this gene;		18108351, 21906765, 21906767, 18108370,
		COMPLEAT 9 X23309:3 Comes from this gene; CDNA EST		18108374, 35696423, 264636, 264639
		elegans]		
2999		94847055 (5997, 5998) Novel Protein sim. GBank	UNCLASSIFIED	56182575 22278996 29147620 29331825
		gij115408jspjP18835jCC19_CAEEL - CUTICLE		29146498 29146499 264905 68712502
		COLLAGEN 19		265006, 265009, 21906754, 85658542.
				18108351, 29148627, 29148629, 60170615.
				33657109, 27486262, 18108370, 18108374,
				264556, 264557, 264558, 60170394,
				18108385, 264563
3000	(0009 '6669' 0266606	95099370 (5999, 6000) Novel Protein sim. GBank gi 1163174 (U32575) - similar to	UNCLASSIFIED	264887, 22278997, 22278999, 264259.
		yeast Sec6p, Swiss-Prot Accession Number P32844, similar		29331822, 29331824, 35696052, 29146498,
		to mammalian B94, Swiss-Prot Accession Number Q03169;		264508, 264905, 264906, 264907, 264908,
		Method: conceptual translation supplied by author [Rattus		264909, 264510, 264511, 265006, 265007,
		norvegicus]	<u> </u>	265008, 265009, 264910, 33657402, 264757,
				264595, 264596, 264758, 21906754, 265011,
				264600, 265017, 265018, 264605, 265019.
				264760, 264761, 264762, 264681, 264682,
			.,	264764, 264288, 264685, 264766, 264686,
		-	<u> </u>	264768, 264769, 21906765, 21906768.
				35695917, 265020, 264691, 264692,
				33657023, 264693, 33657109, 33657182.
				27486261, 264628, 264629, 18108374,
			,	18108376, 35696423, 35695855, 264630,
				264631, 264632, 264634, 264635, 264636,
				264637, 264556, 264638, 264639, 60170394,
			<u> </u>	83373044, 20798451, 22279002, 264563,
2000	00070464 (0004 0000)	North Charles of Contract of C	Ì	264486, 264567
	(znna 'inna) scravinae	ocovious (acovi), ocovis) Novem Profession Simi, Cosan (gildvida / Oktobio/3).	cathepsin	18108394, 52646842, 56182575, 29331824,
		ruddive gene, benscan predictions confirmed by EST	<u> </u>	29331825, 29331827, 264910, 33109954,
	<u></u>	splicing.; coded for by human cDNAs AA122029	<u></u>	52644296, 265017, 265019, 264288, 265020,
		(NID:g1678048), D31562 (NID:g644442), AA158721		265021, 52644150, 284692, 35695763,
	- <del></del>	(NID:g1733515), R59640 (NID:g830335) and F13082	=-	55810764, 35696423, 56182323, 18108387,
5000	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	(NID:g709111) [Homo sapiens]		264563, 264564
	07718157 (6003, 6004)	o/ / 1015/ (auto), but ) Novel Protein sim. GBank gil3599478 (AF085185) - Myosin-	UNCLASSIFIED	264488, 29331824, 29331825, 29331826,
	<del></del> -	IA [Acanthamoeba castellanii]		29331827, 29331828, 264906, 264510,
				265009, 21906754, 264682, 264688,
				33857023, 264565

3003	86648079 (6005, 6006)	3003   86648079 (6005, 6006)   Novel Protein sim. GBank gi 1754969 (U30292) - collagen	Contains protein domain (PF01391) - collagen	collagen	264512, 264593, 264564, 264567, 264486
		type Aill aipna-1 chain [Mus musculus]	Collagen triple helix repeat (20  copies)	_	
304		88066876 (6007, 6008) Novel Protein sim. GBank gi[2224629 dbj BAA20802  -  (AB002342) KIAA0344 [Homo sapiens]			29331830, 21906769, 264691, 33657109, 263972, 18108385
3005		87794843 (6009, 6010) Novel Protein sim. GBank gil4680659 gb AAD27719.1 AF13294 - (AF132944) CGI-10 protein [Homo sapiens]	Contains protein domain (PF01360) - oxygenase Monooxygenase	oxygenase	29331822, 29331824, 29331827, 60433438, 265011, 265019, 21906766, 21906767, 21906768
					60170394, 22279002, 264567
3006		87422224 (6011, 6012) Novel Protein sim. GBank gil3930525 (AF064447) - sex- Idetermination nrotein bomolog Fem1s Mins musculus)	Contains protein domain (PF00023) - MHC	MHC	264259, 29331822, 264512, 21906754,
			Ank repeat		265018, 264687, 21906765, 264691, 264555,   264556, 264558, 18108385
3007		90936005 (6013, 6014) Novel Protein sim. GBank gilz565052 (U80738) - CAGH1a	Contains protein domain (PF00096) - transcriptfactor	transcriptfactor	52644507, 52645156, 65274572, 264909,
		[Homo sapiens]	Zinc finger, C2H2 type		264512, 265018, 264760, 264448, 264765,
					264689, 60170615, 18108374, 20281152,       264636, 52644332
3008	80416249 (6015, 6016)	80416249 (6015, 6016)			264905, 264593, 264766, 264636
3009	(91213387 (6017, 6018)	Novel Protein sim. GBank gij3127193 (AF062389) - kidney-		synthase	52646842, 56182575, 22278995, 22278996,
		specific protein [Rattus norvegicus]	AMP-binding enzyme		264259, 29331825, 29331826, 29331827,
					29331828, 35696052, 264508, 264509,
					264907, 56182435, 264511, 265007, 264512.
					265008, 264757, 264758, 55812038, 264759,
					33109954, 21906754, 265010, 265011,
					264600, 265017, 265018, 265019, 264760,
					18108351, 264288, 264369, 21906764,
					21906/65, 21906/67, 55811957, 265020,
					265021, 264691, 18108368, 27486262,
					20281149, 18108370, 55811576, 264637,
					264556, 264557, 18108381, 264558.
					56182323, 264559, 18108385, 18108388,
3010		95317217 (6019, 6020) Novel Protein sim. GBank	Contains profeio domaio (PE01923)	UNI ACCIEIED	22279UUZ, 284485 284888 284887 21006267 21006769
	•	gil4927370lqblAAD33084_11AF06797 - (AF067972) DNA	Profesio of sokoown function		55811057 2227805 25505017 2228006
		cytosine methyltransferase 3 alpha (Homo sapiens)			22278997 265020 265021 60170615
					264692, 33657023, 29331822, 264693
					18108364, 29331824, 33657109, 60432289.
					29331827, 27486261, 29331828, 264508.
					264909, 55811576, 35695855, 265008,
			-		264556, 60433438, 83373044, 18108387,
					65274727, 60432113, 265017, 22279000,
					265019, 264564, 264682, 264764
רוספ	94323597 (6021, 6022)	94323397 (6021, 6022) Novel Protein sim. GBank	Contains protein domain (PF00153) - transport		35696052, 56182435, 264758, 21906754,
		gijouo2319igapAAD38501.1jAF11883 - (AF118838) citrin;	Mitochondrial carrier proteins		265018, 264760, 264762, 18108351, 264682,
		adult-onset type II cirtuilinemia protein (Homo sapiens)			264448, 21906766, 65274620, 18108374, 1 264482, 264564
3012	87753087 (6023, 6024)			INCLASSIFIED	263972
				1	7,603

			*	*	
ŝ					264558
3019		94148231 (6037, 6038) Novel Protein sim. GBank gij3219332 (AC004020) - Unknown gene product [Homo sapiens]			264569, 52644507, 18108394, 65274572, 56182575, 22278994, 22278995, 56994075, 22278998, 22278995, 56994075, 22278998, 22278999, 264259, 29331822, 29331824, 60432289, 29331827, 264908, 56182435, 265000, 6043229, 265018, 265010, 265000, 6043229, 265018, 265011, 87168559, 265017, 264681, 18108354, 265682, 264682, 264681, 18108354, 264682, 264681, 265691, 33657023, 264692, 264691, 33657109, 52645129, 33657349, 264623, 31857091, 264634, 264633, 318108385, 87168518, 22278000, 22278000, 264683
		943 I 023 I (0039, 0040) Novel Froiein sim. GBank gil34 14609 (AF0b1329) - rjs (Mus musculus)	3414&U9 (AF0b1529) - rjs [Mus Contains protein domain (PF0b415) - ATPase_associated Regulator of chromosome condensation (RCC1)	A i Pase_associated	264488, 263994, 35696286, 264259, 264508, 264909, 264510, 264906, 264907, 264908, 264909, 264909, 264909, 264909, 264909, 264610, 264910, 60174639, 264600, 264603, 264764, 264288, 264369, 26411957, 35695917, 33657023, 264638, 264634, 264635, 264639, 264637, 264639, 264637, 264639, 264639, 264639, 264638, 264567, 264639, 264486
3021		80478512 (6041, 6042) Novel Protein sim. GBank gij3880889jembjCAB09005j - (29559) cDNA EST yk23644.5 comes from this gene; cDNA EST EMBL:C13455 comes from this gene; cDNA EST Yk329g6.5 comes from this gene; cDNA EST CEMSH45R comes from this gene [Caenorhabditis elegans]			264769, 264629, 264482
3022		0			264259, 29331826, 29331828, 264288, 264566
3023		95305484 (6045, 6046) Novel Protein sim. GBank gil416592 sp P32323 AGA1_YEAST - A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR	Contains protein domain (PF00614) - UNCLASSIFIED Phospholipase D. Active site motif		264488, 22278995, 35696286, 22278997, 29331826, 35696052, 264907, 29331830, 52644045, 56182435, 60432229, 264592, 60433356, 60433438, 264689, 21906767, 55811957, 35695917, 265021, 18108376, 263978, 264635, 222279000
3024	_	0		UNCLASSIFIED	60432049, 264760, 21906769, 55811957, 35695917, 264690, 264555, 264559
3025		65706629 (6049, 6050) Novel Protein sim. GBank gil295671 (L11275) - selected as a weak suppressor of a mutant of the subunit AC40 of DNA dependant RNA polymerase I and III [Saccharomyces cerevisiae]			264593, 55811576

	ASSIFIED 18108394, 22278995, 22278996, 35696286, 22278997, 22278998, 60432049, 2564259, 22278998, 60432049, 2564259, 29331822, 29331824, 66714117, 29331825, 60432289, 29331826, 29331827, 29331827, 29331827, 29331827, 256960, 264909, 264907, 29331830, 265009, 264909, 264910, 33657402, 265009, 265009, 264901, 33657402, 265008, 264600, 265011, 87168559, 264600, 265018, 18108351, 264682, 264683, 264689, 21906765, 21906766, 21906767, 21906768, 21906765, 21906766, 21906767, 21906768, 21906769, 29148829, 33657109, 27486261, 18108374, 55811576, 35696423, 6527023, 264599, 264557, 264638, 60170394, 264558, 264557, 264638, 60170394, 264568, 22279000, 22279002		UNCLASSIFIED (22278997, 22278999, 29331827, 264905, 264509, 264509, 264510, 264511, 264512, 87168474, 265019, 18108351, 21906768, 264534, 264690, 264693, 263969, 18108370, 264558, 22279000, 22279002, 264482		UNCLASSIFIED 22278995, 22278997, 22278999, 22278999, 264259, 29331822, 29331824, 29331826, 29331827, 29331828, 35696052, 264908, 265018, 21906765, 21906766, 21906767, 21906768, 265021, 263974, 18108374,
	Contains protein domain (PF01529) - UNCLASSIFIED DHHC zinc finger domain	Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat	חאכר	Contains protein domain (PF00001) - tm7 7 transmembrane receptor ((rhodopsin family)	
3026   87643662 (6051, 6052) Novel Protein sim. GBank gij3024052 sp P97924 KARI_RAT - KALIRIN (PAM COOH-TERMINAL INTERACTOR PROTEIN 10) (P-CIP10)	94844563 (6053, 6054) Novel Protein sim. GBank gil4929647[gb]AAD34084.1 AF 15184 - (AF 151847) CGI-89 protein [Homo sapiens]	94231997 (6055, 6056) Novel Protein sim. GBank gij3080521lemb[CAA18650] - (AL022599) hypothetical protein [Schizosaccharomyces pombe]		87544928 (6059, 6060) Novel Protein sim. GBank gij3757726 emb CAA18782  - (AL022727) dJ80119.1 (olfactory receptor-like protein (hs6M1-1)) [Homo sapiens]	91677953 (6061, 6062) Novel Protein sim. GBank gi 4530587 gb AAD22105.1] - (AF132000) TADA1 protein [Homo sapiens]
1026 87643662 (6051, 6052)					3031   91677953 (6061, 6062)

3032	94130124 (6063, 6064)		Contains protein domain (PF00534) -	synthase	22278996, 35696286, 264259, 29331824,
		M. musculus MER5 and other AHPC/TSA proteins [Caenorhabditis elegans]	Glycosyl transferases group 1		29331828, 264907, 29331830, 264758,
					264288, 21906769, 265021, 264693,
	-				35696423, 35695855, 264636, 56182323,
					83373044, 87168518
3033	95308321 (6065, 6066)	95308321 (6055, 6066) Novel Protein sim. GBank foil50315731effNP 005712 1nACTR - ARP3 (actio-related	Contains protein domain (PF00022) - struct	struct	35696286, 264259, 29331826, 35696052,
		Interest 3, yeast) homolog			264300, 204303, 204300, 264307, 264306. 264909 265008 284591 21906754 265010
					265010 264681 264360 264768 21006764
					203019, 204001, 204309, 204700, 21900704, 0 91006768 35605017 33657033 264628
					21900100, 33093917, 33037023, 204020, 35605855 364632 364635 364630 364483
					264563
3034	80415373 (6067, 6068)			UNCLASSIFIED	264906, 264907, 264510, 264592, 265010,
					264762, 264766, 264637, 264638, 264486
3032		91220692 (6069, 6070) Novel Protein sim. GBank gi[3738207]emb CAA21262  -		UNCLASSIFIED	264636
		(ALU31853) conserved ATP-GTP binding protein [Schizosaccharomyces pombe]			
3036		91718323 (6071, 6072) Novel Protein sim. GBank		kinase	264907, 33657402, 265021
		oil7288371solP391941ALU7 HUMAN - 1111 ALU SUBEAMILY			
		SQ WARNING ENTRY IIII			
3037	95307434 (6073, 6074)	95307434 (6073, 6074) Novel Protein sim, GBank gil4406590lgblAAD200401.			265017
		(AF131766) Similar to Ena-VASP like protein [Homo			
		sapiens]			
3038	85421807 (6075, 6076)	3038   95421807 (6075, 6076) Novel Protein sim. GBank	Contains protein domain (PF00627) - UNCLASSIFIED	Γ	22278996, 22278997, 264259, 264905,
		gi 5360093 gb AAD42865.1 AF15509 • (AF155099) NY-REN UBA domain	UBA domain		265007, 265009, 60433356, 21906754,
		18 antigen [Homo sapiens]			265018, 265019, 18108351, 264687.
					21906765, 265020, 265021, 65274620,
					27486262, 264636, 56182323, 18108385,
2020	10200 4200 73000000	The second of th		1	222/9000
	01335631 (0011, 0010)	07.332237 (0077, 0070) (NOVELPTOTEIN SIM. GBANK 91/4757128/emb CAB42094.1 -		UNCLASSIFIED	35696286, 29331828, 264109, 264110,
		לארכיסי בין לאבן לא הספונו (עשוות שומה המוכח)			264311, 265007, 21906734, 265011, 264681,
					254583, 254587, 21905/58, 254591, 18108370, 263572, 264620, 18108374
					263977, 35696423, 264564, 18108391
3040	90933517 (6079, 6080)	90933517 (6079, 6080) Novel Protein sim. GBank gil4884278 emb CAB43247.1  -			264692, 264558, 18108382, 18108385,
		(AL050037) hypothetical protein [Homo sapiens]			264567
304	88312357 (6081, 6082)	Novel Protein sim. GBank gil3876073 emb CAB04122.1  -		UNCLASSIFIED	56994075, 22278997, 22278998, 29331827,
		(Z81505) similar to Zinc finger, C3HC4 type (RING finger);			33656970, 33109954, 21906754, 87168559,
		cDNA EST EMBL: D28025 comes from this gene; cDNA			264600, 264683, 21906765, 21906768,
		EST EMBL: D28024 comes from this gene; cDNA EST			22279002
		EMBL: D33210 comes from this gene; cDNA EST			
,	10000 00000	PETAGOGO COSTANTINE COMES TROM THIS			
3042	85/49402 (6083, 5084)	Novel Protein sim. GBank gi[790236 (U21156) - sarcolemmal associated protein-2 (Ovctolanus cuniculus)		glycoprotein	264636
		Company on Spring Company			

3043	87773026 (6085, 608)	3043   87773026 (6085, 6086) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	35696286, 60424269, 35696052, 264508, 264905, 66712502, 56182435, 55811386, 52644296, 55811150, 35695917, 60170615, 33657109, 18108374, 264634, 60431850
3044		87646182 (6087, 6088) Novel Protein sim. GBank gil4104922 (AF042276) - 0251 homolog [Pseudomonas pulida]	Contains protein domain (PF01209) - glycoprotein ubiE/COQ5 methyltransferase family	glycoprotein	22278996, 22278998, 22278999, 29331824, 56182435, 264511, 265007, 60170831, 60432229, 60433356, 33109954, 18108351, 264288, 35695917, 18108368, 18108370, 60170394
3045		94127598 (6089, 6090) Novel Protein sim. GBank gil4589680 dbj BAA76859.1   - (AB023232) KIAA1015 protein [Homo sapiens]	Contains protein domain (PF00096) - dna_rna_bind Zinc finger, C2H2 type	dna_ma_bind	264488, 264259, 35696052, 264508, 264905, 264509, 264509, 264509, 264906, 264907, 264909, 264511, 265006, 264591, 284593, 33109954, 264604, 264768, 24906765, 21906768, 55811957, 35695917, 27486262, 18108370, 264628, 18108374, 264566, 264566
3046				UNCLASSIFIED	22278999, 29331822, 29331824, 29331825, 29331826, 60432289, 29331827, 29331828, 264906, 52646317, 55811957, 60432113, 22279000, 22279002, 264482, 264564
3047				UNCLASSIFIED	264468, 22278996, 22278997, 22278999, 29331824, 29331825, 56182435, 264511, 265008, 265011, 265017, 264766, 21906768, 21906769, 35695917, 52644150, 33657349, 65274791, 35695855, 264555, 60432113, 22279000, 264566
3048		87629419 (6095, 6096) Novel Protein sim. GBank gi 4588034 gb AAD25962.1 AF09287 - (AF092878) zinc RtNG finger protein SAG (Homo sapiens)	Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)	UNCLASSIFIED	264102, 29148784
3049		) Novel Protein sim. GBank gij5454158jrefNP_006286.1jpVARS - valyl-tRNA synthetase 1	Contains protein domain (PF01406) - UNCLASSIFIED IRNA synthetases class I (C)	UNCLASSIFIED	22278997, 29331826, 264907, 264758, 87168559, 265018, 264448, 21906766, 265020, 33657109, 35695855, 60432113, 22279000
3050		87643679 (6099, 6100) Novel Protein sim. GBank gil4589642 dbj BAA76843.1  - (AB023216) KIAA0999 protein [Homo sapiens]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264259, 29331825, 264909, 265007, 264512, 265019, 264288, 21906766, 265020, 264693, 18108385, 56526486, 87168518, 22279002, 264566
3051		(6			22278997, 264595, 265019, 264288, 264693, 87168518
3052		57108030 (6103, 6104) Novel Protein sim. GBank gij117528jspjP14755jCRYL_RABIT - LAMBDA- CRYSTALLIN		dehydrogenase	264534

65274572, 56181686, 22278995, 35696286, 22278998, 264259, 60432289, 265008, 265009, 60433438, 21906754, 265010, 87168559, 264603, 265018, 265019, 264763, 264764, 264288, 21906765, 21906766, 21906768, 21906768, 35695917, 18108374, 35696423, 264638, 56182323, 22279000, 264563	35696286, 35696052, 29331830, 264908. 264909, 264512, 284910, 265017, 264604. 264766, 265020, 33657109, 264628. 35695855, 264636, 264566, 264486.	60424179, 65274572, 56182575, 35696286, 22278996, 22278999, 60432049, 264259, 60424269, 60432289, 35696052, 56182435, 265006, 265009, 60170831, 60433229, 265009, 6013356, 264594, 60433438, 21906754, 55811386, 265011, 87168559, 265019, 18108351, 264683, 264288, 264369, 264689, 21306768, 55811957, 35695917, 60170815, 33657023, 5527462109, 35695763, 60431528, 18108374, 55810764, 55811576, 35696423, 56182323, 60170394, 18108385, 60432113, 264564, 264565,	264488, 264569, 18108394, 52646842, 22276897, 22278998, 22278999, 264259, 66714117, 29331826, 29331827, 35696052, 264308, 264509, 264905, 264906, 264906, 264906, 264906, 264906, 265009, 264906, 264907, 265009, 264909, 264910, 33657402, 265017, 265019, 264760, 18108351, 264761, 265017, 265019, 264760, 18108351, 264761, 264763, 264764, 264289, 264689, 21906765, 21906766, 21906767, 21906765, 21906766, 21906767, 265021, 264629, 264639, 336557109, 18108370, 264629, 264634, 264639, 264634, 264639, 264631, 264634, 264634, 264636, 264564, 264566, 264486, 264567, 264567, 264566, 264486, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264536, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 26457, 264567, 264567, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457,
UNCLASSIFIED	UNCLASSIFIED	transport	glycoprotein
			·
Novel Protein sim. GBank gil3947613 emb CA419465.1  - (AL023828) cDNA EST EMBL:M89008 comes from this gene; cDNA EST yk282d3.5 comes from this gene [Caenorhabditis elegans]	86943510 (6107, 6108) Novel Protein sim. GBank gij1076211 pirj S50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardtii	Novel Protein sim. GBank gild680655[gb AAD27717.1 AF13294 - (AF132942) CGI-08 protein [Homo sapiens]	91661636 (6111, 6112) Novet Protein sim. GBank gij728837jspjP39194/ALU7_HUMAN - !!!! ALU SUBFAMIL Y SQ WARNING ENTRY !!!!
3053   95350373 (6105, 6106) Novel Protein sim. GBank gil (AL023828) cDNA EST EMB gene; cDNA EST yk282d3.5 [Caenorhabditis elegans]	3054 86943510 (6107, 6108) N	3055   95350537 (6109, 6110)   9	3056 91661636 (6111, 6112) (

264508, 264905, 264907, 264908, 264909, 264510, 264512, 264910, 264592, 264594, 264767, 18108374, 264635, 264555, 26486 264639, 264563, 264563, 264563, 26486	264693	22278995, 22278996, 22278997, 22278998, 22278999, 264490, 264259, 29331824, 29331825, 29331827, 35696052, 29331828, 265007, 60433438, 265017, 265018, 265019, 264681, 264448, 264288, 264768, 21906765, 21906767, 21906769, 29148784, 265022, 52644150, 18108370, 264636, 18108355, 264563, 264567	264567	264112, 52644296, 21906768, 33657023, 263974, 18108385	264908, 265008, 18108351, 264566	18108359, 264558	52646365, 52646842, 65274572, 56182575, 52646365, 52646842, 65274575, 56182575, 56181686, 22278996, 22278996, 22278997, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 264908, 66714117, 264508, 264907, 264908, 2644296, 265009, 60432229, 60433438, 55812038, 2644296, 2650018, 264697, 264689, 21906768, 264691, 264697, 264689, 18108370, 18108377, 55811576, 264636, 56182323, 264589, 264693, 26279000, 22278998, 264093, 264094, 264995, 29331824, 60424269, 66714117, 264096, 29331824, 60424269, 66714117, 264096, 29331824, 60424269, 66714117, 264593, 265011, 265019, 18108351, 264766, 264593, 26575007, 2932007, 264593, 2047806, 2932007, 2047807, 2644893, 204780769, 2932007, 264593, 264466, 2932007, 264593, 264466, 29320007, 29320007, 264482	264566, 264567
struct	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transferase	UNCLASSIFIED	UNCLASSIFIED	
		Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)					Contains protein domain (PF00403) - UNCLASSIFIED Heavy-metal-associated domain UNCLASSIFIED	
95412746 (6113, 6114) Novel Protein sim. GBank gil3878119 emb CAA88880  - (Z49068) similar to GTP-binding protein; cDNA EST EMBL:M89111 comes from this gene; cDNA EST EMBL:D27709 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D73788 comes from this gene; cDNA EST		87829425 (6117, 6118) Novel Protein sim. GBank gil4588034[gb]AAD25962.1[AF09287 - (AF092878) zinc RING finger protein SAG [Homo sapiens]			87619465 (6123, 6124) Novel Protein sim. GBank gil4454690lgblAAD20963  - (AF070857) glutathione S-transferase subunit 13 homolog [Homo sapiens]	3063 80078023 (6125, 6126) Novel Prolein sim. GBank gi[2246532 (U93872) - ORF 73, contains large complex repeat CR 73 (Kaposi's sarcoma-associated hemosvirus)	91241526 (6127, 6128) Novel Protein sim. GBank gil4240315 dbj BAA74936.1  - (AB020720) KIAA0913 protein [Homo sapiens] 91639201 (6129, 6130) Novel Protein sim. GBank gil5656743 gb AAD45960.1 AC00506 - (AC005067) Supported by Human EST H08032.1 (NID:g872854), mouse EST AA870042.1 (NID:g2965487), and genscan [Homo sapiens]	
95412746 (6113, 6114)	79646226 (6115, 6116)	87629425 (6117, 6118)	79346691 (6119, 6120)		87619465 (6123, 6124)	80078023 (6125, 6126)	91241526 (6127, 6128)	
3057	88	3059	3060	3061	3062	306	3065	

18108397, 22278995, 56994075, 22278996, 264905, 66712502, 265006, 264512, 264910, 264758, 60174639, 264760, 18108351, 264764, 264683, 18108359, 264692, 18108364, 18108379, 264567	264488, 264489, 35696286, 22278996, 56994075, 264259, 29331822, 29331825, 35696052, 29331822, 29331825, 26594075, 264509, 264509, 264509, 264510, 264511, 264512, 265008, 264509, 264511, 264511, 264512, 265008, 264509, 264511, 264511, 264512, 265008, 264509, 264501, 264501, 264501, 264601, 264601, 264602, 264603, 264604, 264605, 265019, 264760, 264369, 264760, 264760, 264369, 264601, 264603, 264603, 264601, 264603, 264603, 264601, 264603, 264601, 264603, 264603, 264603, 264603, 264603, 264603, 264603, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264563, 264565, 264566, 264567, 264483, 26279000, 22279002, 264563, 264483, 262799000, 22279002, 264567, 264483, 262799000, 22279002, 264567, 264486, 264564, 264565, 264566, 264567, 264567, 264486,	264112	22278996, 56994075, 22278998, 22278999, 2264259, 264107, 264905, 29331630, 52644045, 264107, 264905, 29331630, 264594, 33657402, 21906754, 33109954, 87168474, 87168559, 265017, 264448, 264764, 264683, 264766, 52644229, 21906765, 21906766, 21906768, 21906769, 60170615, 33657023, 18108370, 18108336, 264634, 264557, 60170394, 56182323, 18108385, 87168518, 22279000, 264482
UNCLASSIFIED	struct		struct
	Contains protein domain (PF00787) - struct		Contains protein domain (PF01926) - struct GTPase of unknown function
3066   91224437 (6131, 6132) Novel Protein sim. GBank gil4884268 emb CAB43245.1  - (AL050028) hypothetical protein [Homo sapiens]	95422551 (6133, 6134) Novel Protein sim. GBank gil4689258[gb]AAD27832.1 AF12185 - (AF121859) sorting nexin 9 [Homo sapiens]	35, 6136)	95412753 (6137, 6138) Novel Protein sim. GBank gij3878119jembjCAA88860j - (Z49068) similar to GTP-binding protein; cDNA EST EMBL:M89111 comes from this gene; cDNA EST EMBL:D27709 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D73788 comes from this gene; cDNA EST
91224437 (613	3067 95422551 (6133	3068 85360651 (6135, 6136)	3069 95412753 (613)

264488, 22278994, 22278995, 22278996, 56994075, 22278997, 22278999, 264259, 29331822, 29147620, 29331824, 66714117, 29331826, 29146498, 29146499, 66712502, 29331836, 52644045, 56182435, 264511, 265007, 264512, 264910, 60170831, 264511, 265019, 18108351, 264448, 264683, 264288, 21906764, 21906765, 21906766, 21906766, 21906767, 21906769, 25644150, 264691, 265021, 60170615, 52644150, 264628, 18108370, 18108381, 60170394, 56182323, 264568, 264563, 18108387, 56526486, 264563, 18108387, 56526486, 264563, 18108387, 56526486, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264566, 264563, 264566, 264563, 264566, 264563, 264566, 264563, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264566, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 26466, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 26466, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 264666, 2646	264488, 65274572, 18108398, 22278996, 35696286, 22278997, 22278998, 22278999, 264259, 29331822, 65714117, 29331826, 264259, 29331828, 29146499, 264907, 264908, 29331830, 264909, 265900, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265102, 265117, 26502, 264691, 265117, 26502, 264691, 2651176, 2551176, 26502, 264691, 2651176, 2651176, 26502, 264691, 2651176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551176, 2551177, 26553, 26557, 60170394, 83373044, 60432113, 22279000, 22279002, 264563, 264565, 264565, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567	56182575, 29331822, 29331824, 29331825, 29146498, 264908, 52644045, 56182435, 265009, 60433438, 55812038, 18108351, 264683, 264369, 52644229, 52644150, 33657023, 264693, 33657109, 18108374, 55811576, 65274791, 264555, 56182323, 60432113, 264564
synthase	tgf	transcriptfactor
,	Contains protein domain (PF00085) - tgf Thioredoxin	
Novel Protein sim. GBank gij3877788 emb CAB05527  - (283110) cDNA EST yk472b5.3 comes from this gene; cDNA EST yk472b5.3 comes from this gene; cDNA EST yk472b5.5 comes from this gene; cDNA EST yk468c10.3 comes from this gene; cDNA EST yk468c10.5 comes from this gene; cDNA EST yk468c10.5 comes from this gene; cDNA EST was a come from this gene; cDNA EST EM	94325573 (6141, 6142) Novel Protein sim. GBank gil4502425 ref NP_001709.1 pBMP6 - bone morphogenetic protein 6 precursor	95115892 (6143, 6144) Novel Protein sim. GBank gi[1263289 (U47856) - fibroin-4 [Araneus diadematus]
3070   94319173 (6139, 6140)   Novel Protein sim. GBank gi (283110) cDNA EST yk474a7 3 come. cDNA EST yk474a7 3 come. yk472b5.5 comes from this gene; cDNA this gene; cDNA EST EM	3071 94325573 (6141, 6142) N	3072   95115892 (6143, 6144) [[]

2072	106447740 /6446 6446	2072   06447740 /6448   6446)   11   11   11   11   11   11   11			
3		NOVEL FIOLEIN SIIII, ODGIIN 9i 134840 sp P22528 CORB_HUMAN - CORNIFIN B (SMALL PROLINE-RICH PROTEIN IB) (SPR-IB) (14.9 KD PANCORNULIN)		UNCLASSIFIED	264 / 69
3074	88089351 (6147, 6148)	88089351 (6147, 6148) Novel Protein sim. GBank gil3419847 (AC004982) - similar to yeast hypothetical protein ybk4; similar to P38164		UNCLASSIFIED	264488, 265019, 264448, 264288, 21906767, 264693, 18108368, 18108370, 18108374,
3075		88095752 (6149, 6150) Novel Protein sim. GBank gil465749fjreff)P_000456.1[pBARD - BRCA1 associated RMG domain 1	Contains protein domain (PF00023) - homeobox Ank repeat	<b>homeobox</b>	264509, 264907, 264689, 264693, 56526486
3076	87819219 (6151, 6152)			UNCLASSIFIED	18108398, 29331822, 29331827, 60432229, 265017, 264691, 264693
3077		88734277 (6153, 6154) Novel Protein sim. GBank gij3023956 sp Q00808 HET1_PODAN - VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat	kinase	65274572, 35696052, 264511, 60170831, 87168474, 264369, 35695917, 33657182, 27486264, 33657349, 35695763, 35695855, 264339
3078		88089355 (6155, 6156) Novel Protein sim. GBank gil3900850 (AC004994) - similar to d1026456 (PID:g3043724) [Homo sapiens]			22279002
3079		87821893 (6157, 6158) Novel Protein sim. GBank gij3875410 emb CAB02876  - (281052) Similarity to Yeast ABC1P protein (SW-ABC1_YEAST); cDNA EST yk229g8.3 comes from this gene; cDNA EST yk229g8.5 comes from this gene [Caenorhabditis elegans]		transport	29331824, 29331826, 264758, 55811386, 265017, 55811150, 52644229, 21906768, 265020, 265021, 264693, 18108376, 264631, 52644332, 22278002
3081		98094864 (6161, 6162) Novel Protein sim. GBank J WARNING ENTRY !!!!		UNCLASSIFIED	264490, 264259, 29331824, 66714117, 264490, 264259, 29331824, 66714117, 29331825, 29331827, 29331827, 29331828, 29331827, 29331827, 29331828, 29331827, 29331827, 29331828, 264699, 264508, 264508, 264608, 264683, 264763, 264683, 264764, 264288, 264684, 264268, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264690, 264692, 33657023, 264693, 65274620, 33657023, 264693, 65274620, 3365709, 27488261, 35695763, 264689, 264909, 264510, 264589, 264909, 264510, 264511, 265009, 264760, 264511, 265009, 264690, 264631, 264683, 264631, 264683, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264633, 264655, 264633, 264655, 264633, 264655, 264633, 264655, 264633, 264655, 264633, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264656, 264666, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26
3082	3082  80310121 (6163, 6164)				264764, 55811957, 264555, 264564

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UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	dehydrogenase
						Contains protein domain (PF00725) - dehydrogenase 3-hydroxyacyl-CoA dehydrogenase
3083   88095756 (6165, 6166)   Novel Protein sim. GBank gil888241 (U29488) - C56C10.3 gene product [Caenorhabditis elegans]	87448568 (6167, 6168) Novel Protein sim. GBank gil476774 pir  A37475 - probable structural component p38 - borna disease virus	87795781 (6169, 6170) Novel Protein sim. GBank gi 2565057 (U80741) - CAGH44 [Homo sapiens]	87769942 (6171, 6172) Novel Protein sim. GBank gil3894189 (AC005662) - hypothetical protein [Arabidopsis thaliana]		91224441 (6175, 6176) Novel Protein sim. GBank gil335304 (AF001549) - Unknown gene product [Homo sapiens]	95361242 (6177, 6178) Novel Protein sim. GBank gil4689146[gb]AAD27782.1 AF07704 - (AF077049) lambda- 3-hydroxyacyl-CoA dehydrogenase crystallin [Homo sapiens]
88095756 (6165, 6166)	87448568 (6167, 6168)			87462988 (6173, 6174)	91224441 (6175, 6176)	95361242 (6177, 6178) '
3083	3084	3085	3086	3087	3088	3089

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UNCLASSIFIED	UNCLASSIFIED
Novel Protein sim. GBank gi[1354050 (U47024) - MEM3 [Mus musculus]	95317424 (6181, 6182) Novel Protein sim. GBank gij3873932 emb CAB01859  - (Z79596) Simialrity to Bovine aspartyl beta hydroxylase (TR:G162694); cDNA EST EMBL:D27916 comes from this gene; cDNA EST EMBL:D27915 comes from this gene; cDNA EST EMBL:D64881 comes from this gene; cDNA EST EMBL:D64881 comes from this gene; cDNA EST EMBL:D68139 comes from this gene; cDNA
3090 95342371 (6179, 6180) Novel Protein sim. GBank [Mus musculus]	3091 95317424 (6181, 6182)  N

5314592 (6183, 6184)	3092   95314592 (6183 6184) Novel Protein sim GBank	Contains protein domain (PF00333) - Iribosomalprot	bosomalprot	264488, 60424179, 18108396, 22278995,
	gil17107561spiP15880IRS2 HUMAN - 40S RIBOSOMAL	Ribosomal protein S5		56994075, 22278996, 35696286, 22278997,
	PROTEIN S2 (S4) (LLREP3 PROTEIN)			22278998, 60432049, 264259, 29331822,
				29331824, 29331825, 29331826, 29331827.
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8457 (6185, 6186	94318457 (6185, 6186) Novel Protein sim. GBank gil5002587 emblCAB44347.11 -	1	UNCLASSIFIED	264259, 29331824, 35696052, 264905,
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				264369, 264288, 264766, 21906767,
				35696423, 83373044, 18106363
6675 (6187, 6188	94316675 (6187, 6188) Novel Protein sim. GBank gi 400734 sp  931044 PBP_RAT	400734 sp P31044 PBP_RAT -  Contains protein domain (PFU1161) -  Collagen	collagen	10100396, 204239, 00432269, 29331047.
	PHOSPHATIOYLE I HANOLAMINE-BINDING PROTEIN (23 Prosphatioyletnanoiamine-binding)	Prosphatidyletnanolamine-billionig		264501 264603 65274701 56182323
	KD MORPHINE-BINDING PROTEIN) (P23K)	protein		204031, 204033, 03214131, 33102323,
				204304, 204303

94648162 (6189, 6190) Novel Protein sim. GBank gij3882221 [dbj]BAA34470.1] - (AB018293) KIAA0750 protein [Homo sapiens] (AB018293) KIAA0750 protein [Homo sapiens] (AB018293) KIAA0750 protein [Homo sapiens] (A22395) KIAA0750 protein [Homo sapiens] (A22395) Movel Protein sim. GBank gij4468288 emb]CAB37981] - (A22395) Movel Protein sim. GBank gij4468288 emb]CAB37981] - (A1022395) Movel Protein sim. GBank gij4468288 emb]CAB37981] - (A102295) GBOOK GB	Contains protein domain (PF01454) - UNCLASSIFIED  - Contains protein domain (PF00307) - struct Calponin homology (CH) domain  - Contains protein domain (PF00646) - UNCLASSIFIED  EST F-box domain.	UNCLASSIFIED UNCLASSIFIED cyto450	18108397, 56182575, 22278995, 35696286, 56994075, 22278997, 22278999, 264259, 60432049, 66714117, 29331825, 60432289, 35696052, 33656970, 29146499, 264508, 264605, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264501, 265010, 265011, 265012, 265019, 264760, 265011, 265012, 264760, 265011, 264063, 264760, 264018, 264681, 264682, 264682, 264760, 265019, 264760, 264019, 264760, 264019, 264760, 264019, 264019, 264019, 264760, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 264019, 26
87602421 (6199, 6200) Novel Protein sim. GBank gi 1083764 pir  B48013 - proline- rich proteoglycan 2 precursor, parotid - rat	line-	UNCLASSIFIED	56182323, 22279002, 264563 29331825, 60432289, 35696052, 264910, 60432229, 264592, 264288, 264693, 263967,
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3102	91220892 (6203, 6204)	3102  91220892 (6203, 6204) Novel Protein sim. GBank	Contains protein domain (PF00018) - Istruct	struct	35696286 22278996 22278999 29331827
		gij5305706 gb AAD41781.1 AF12853 - (AF128536)	SH3 domain		35696052, 264909, 264512, 265008,
		cytoplasmic phosphoprotein PACSINZ [Homo sapiens]			60170831, 60433356, 33109954, 18108351,
					264684, 264689, 21906767, 60170615,
					264692, 33657023, 264638, 22279000,
2463	_	00000004 (CODE CODE) N (C			264482, 264564
3		NOVEI Frotein sim. CBBNX gij46645ppp35292jRB17_MOUSE - RAS-RELATED		UNCLASSIFIED	35695917, 264565
240	_	87340632 (6207 6208) Navel Bestein Charle			
<u> </u>		Novel Protein Sim. Gbank		UNCLASSIFIED	264259, 264684, 264532, 33657182, 264558
		STF CDNA 6		4	
3105	94148603 (6209, 6210)				22278997, 264259, 29331824, 35696052.
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					264594, 264758, 264760, 264681, 264762,
					264764, 264288, 264768, 264768, 264687,
					264769, 21906766, 21906768, 35695917,
					33657023, 264692, 264693, 264628, 264629.
					35695855, 264630, 264631, 264632, 264634.
					264635, 264637, 264638, 264639, 83373044,
					264404 22279002 264563 264565 264566
					264486, 264567 264486, 264567
3106		95361416 (6211, 6212) Novel Protein sim. GBank gi 1938574 (U97190) - B0025.2			22278996, 22278997, 22278998, 22278999.
		gene product [Caenorhabditis elegans]			264092, 264093, 264094, 29331822, 264906.
		•			264907, 264908, 52644045, 56182435.
					264112, 265008, 265009, 55812038, 265017.
			٠		265018 264683 264686 264687 264768
					52644229 21906765 21906768 21906769
					55811957, 265020, 265022, 264690.
					52644150, 264692, 264693, 18108370.
					18108377, 55811576, 56182323, 18108385.
					18108388, 22279000, 264563
3107		95343272 (6213, 6214) Novel Protein sim. GBank gij3341441 emb CAA76851  -			22278995, 22278996, 35696286, 22278997,
		(Y17794) winged-helix transcription factor [Gallus gallus]			22278999, 264091, 264093, 264259,
					29331822, 29331825, 29331826, 60432289,
					29331827, 29331828, 33656970, 264105,
_					264512, 265009, 60433356, 60433438,
					265011, 265017, 265018, 21906765.
					21906766, 21906767, 21906769, 265021,
					264691, 33657109, 27486261, 27486265,
					18108370, 263972, 18108374, 55811576,
					18108385, 56526486, 264482, 264487
3108	87340635 (6215, 6216)	87340635 (6215, 6216) Novel Protein sim. GBank		UNCLASSIFIED	56182435, 264288, 264690, 264564
		griposzzov jretjink judogao. Ijp i ooc - tumor-suppressing ISTE cona 6			
			T		

3109	94318461 (6217, 621)	3109  94318461 (6217, 6218)   Novel Protein sim. GBank gi 5002587   emb CAB44347.1  -	Contains protein domain (PF00096) - struct	struct	264490, 264908, 265007, 264910, 264593,
		(Y17454) LSFR1 protein [Homo sapiens]	Zinc finger, C2H2 type		264683, 264684, 264687, 21906767,
					21906768, 264693, 18108370, 264629,
					18108374, 264632, 264638, 22279000
3110		=		UNCLASSIFIED	264488, 65274572, 22278995, 22278997,
		hypothetical protein VSP-3 - Chlamydomonas reinhardtii			60432049, 264259, 29331822, 29331824,
					29331825, 60432289, 29331826, 29331827,
					29331828, 264906, 264510, 265006, 265007,
					265008, 265009, 60432229, 33657402.
					60433356, 265011, 87168559, 264600,
					265017, 265018, 265019, 18108351, 264288,
					264369, 21906766, 21906767, 21906768,
					265020, 60170615, 264693, 65274620,
					18108370, 264639, 18108384, 22279000,
			,		264563, 18108390
3111	_	87754512 (6221, 6222) Novel Protein sim. GBank gij3282231 (U75454) - C2H2 type	3282231 (U75454) - C2H2 type Contains protein domain (PF00096) - Itranscriptfactor	transcriptfactor	264488, 18108398, 66712502, 265017,
		zinc finger protein [Homo sapiens]	Zinc finger, C2H2 type		265018, 265019, 264448, 21906767, 265020.
					33657023, 18108365, 18108368, 35696423,
	_				52644332, 18108385, 18108388
3112	_	88043639 (6223, 6224) Novel Protein sim. GBank gij3900848 (AC005023) - match	Contains protein domain (PF00046) - homeobox	homeobox	
		to EST AA361117 (NID:g2013436) [Homo sapiens]	Homeobox domain		
3113	_	88207098 (6225, 6226) Novel Protein sim. GBank gi 2459910 (AF005856) -		tm?	18108397, 22278999, 264259, 29331824,
		anon2A5 (Drosophila yakuba)			35696052, 264907, 264757, 60433438,
					87168559, 264763, 264448, 18108354,
					264288, 21906767, 21906769, 35695917,
					264690, 264691, 264692, 264693, 18108365,
					18108381, 18108384, 18108385, 18108388,
					87168518, 22279000, 22279002
3114	_	79843167 (6227, 6228) Novel Protein sim. GBank gil4966270 gb AAB52261.2  -	Contains protein domain (PF00702) - hydrolase	hydrolase	264909, 56182435, 264910, 21906754
		(U97002) similar to acyl-CoA dehydrogenases and epoxide	_		
		hydrolases; Pfam domain PF00441 (Acyl-CoA_dh).	hydrolase		
		Score=57.4, E-value=1.7e-16, N=2; contains similarity to			
		Pfam domain PF00702 (Hydrolase), Score=57.4, E-			
	_	value=1e-13, N=1 [C			PROFESSION CARROLL CAR
3115	_			transcriptiactor	60424179, 56182575, 264259, 29331824,
_		gij5032225jretjNP_005676.1 pWBSC - Williams-Beuren			60424269, 29331828, 66712502, 264510,
		syndrome chromosome region 11			265007, 60431735, 60433356, 55812038,
					55811386, 265019, 264288, 264689,
					21906769, 264691, 33657023, 264693,
					60431528, 263974, 60431850, 56182323.
					264559, 22279000, 22279002
3116		2)		UNCLASSIFIED	264905, 264758, 21906764, 264690
3117		4)		UNCLASSIFIED	264510, 265011, 18108351, 264288, 264689,
					264691, 18108368, 18108372, 263981,
					264558, 264564

52645156, 52646842, 65274572, 56182575, 22278995, 56994075, 22278996, 35696286, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 66714117, 29331826, 29331827, 3569052, 29331828, 264905, 264510, 284511, 265007, 265009, 264570, 2646317, 21906754, 33657084, 52644298, 87168474, 87168559, 265017, 265018, 264682, 264694, 2644150, 264681, 265020, 265022, 264690, 52644150, 264628, 19006766, 21906768, 21906769, 33657023, 264628, 18108370, 60431528, 18108374, 35696423, 65274791, 60170394, 83746823, 364760, 264628, 18108370, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 2047600, 20476000, 20476000, 20476000, 20476000, 20476000, 20476000, 20476000, 20476000, 2047600000000000000000000000000000000000	265006, 264288	.D 264488, 264509, 264510, 264511, 264512, 264288, 264486	52644507, 52645156, 52646365, 52646842, 22278994, 56994075, 22278996, 22278999, 264259, 29331824, 29331827, 35696052, 52644045, 265008, 52646317, 87168474, 87168559, 21906765, 52644150, 33657023, 18108374, 264637	T		Γ	
J61		UNCLASSIFIED		UNCI ASSIFIED	kirase	UNCLASSIFIED	dehydrogenase
Contains protein domain (PF00008) - 1gf	Contains protein domain (PF00328) - Histidine acid phosphatase				[2439517 (AC002563) - putative Contains protein domain (PF00780) - kinase 15% similarity to P49205 CNH domain ans]		Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase
3118 94655848 (6235, 6236) Novel Protein sim. GBank gij3880563jemb[CAB01444.1[- (Z78016) predicted using Genefinder; similar to serine/fhreonine kinase; cDNA EST yk353d10.5 comes from this gene [Caenorhabditis elegans]		87344040 (6239, 6240) Novel Protein sim. GBank gil5019819[gb]AAD37863.1]AF14315 - (AF143152) putative NADH oxidoreductase complex I subunit [Caenorhabditis letegans]	Novel Protein sim. GBank gi 4501877 ref NP_001088.1 pACR  - acrosin		88083003 (6245, 6246) Novel Protein sim. GBank gi 2439517 (AC002563) - putative RHO/RAC effector protein; 95% similarity to P49205 (PID:g1345860) [Homo sapiens]		91216607 (6249, 6250) Novel Protein sim. GBank gil4980826[gb]AAD35412.1]AE00171 - (AE001714) oxidoreductase, short chain dehydrogenase/reductase family [Thermotoga maritima]
94665848 (6235, 6236)	85728796 (6237, 6238)	87344040 (6239, 6240)	94110735 (6241, 6242)		88063003 (6245, 6246)		91216607 (6249, 6250)
3118	3119	3120	3121	3122	3123	3124	3125

3126	3126  95337205 (6251, 6252)			UNCLASSIFIED	22278999 264490 264259 60432049
					29331822, 60432289, 29146498, 52644045
					56182435 265009 60433438 265010
					87168559 265017 265018 55811150
					264763 264683 264369 264685 29148629
					33657023, 264693, 33657109, 18108374
					55811576, 18108385, 60432113, 22279002
3127		91639233 (6253, 6254) Novel Protein sim. GBank gi 2828280 emb CAA16694.1  -			35696286, 22278996, 22278999, 29331826,
		(AL021687) putative protein [Arabidopsis thaliana]			264908, 60433438, 87168559, 264604,
					21906765, 21906769, 33657023, 33657349,
					264629, 18108374, 18108377, 22279000,
					22279002
3128	87674330 (6255, 6256)	3Bank gij3885828 (AF090133) - lin-7-A	_	misc_channel	22278996, 264259, 52644045, 265008,
		[Rattus norvegicus]	PDZ domain (Also known as DHR or		21906754, 265017, 265018, 21906768,
			GLGF).		18108376, 18108387, 22279000, 22279002
3129	87755412 (6257, 6258)	3129 (87755412 (6257, 6258) Novel Protein sim. GBank gij3135273 (AC003058) -	Contains protein domain (PF00400) - kinase	kinase	56182575, 264259, 29331825, 29331828,
		hypothetical protein [Arabidopsis thaliana]	WD domain, G-beta repeat		52644045, 56182435, 60433356, 264600,
_					264682, 264763, 264764, 264369, 264288,
					264686, 55811957, 264692, 33657023,
					33657109, 60432113, 264564, 264566
3130	14993860 (6259, 6260)	14993960 (6259, 6260) Novel Protein sim. GBank gij3329465 (AF064553) - NSD1 protein IMus musculust			264636
3131	95351469 (6261, 6262)	95351469 (6261, 6262) Novel Protein sim. GBank gi[1848277 (U86136) -	Contains protein domain (PF00400) - UNCLASSIFIED	UNCLASSIFIED	56182575 264259 29331824 264907
		telomerase-associated protein TP-1 [Homo sapiens]	WD domain, G-beta repeat		56182435, 264594, 60433438, 55812038,
					33109954, 21906754, 33657084, 87168474,
					264448, 264786, 21906769, 55811957,
_					265020, 265021, 265022, 60170615,
					33657023, 33657109, 33657182, 27486261,
					33657349, 65274791, 60170394, 56182323.
					83373044, 87168518, 264564

3132	95415459 (6263, 6264)	3132 95415459 (6263, 6264) Novel Protein sim. GBank gil4680647 gblAAD27713.1 AF13293 - (AF132938) CGI-03 protein [Homo sapiens]	Contains protein domain (PF00789) - lubiquitin UBX domain	ubiquitin	52644507, 52646842, 52646365, 65274572, 56182576, 22278994, 22278995, 25586886, 56984075, 22278995, 22278995, 22278996, 22278998, 22278998, 22278998, 22278998, 22278998, 22278998, 2223899, 22331822, 29331824, 29331826, 2642080, 29331822, 29331827, 29331828, 264910, 6017038, 5264045, 56182435, 264910, 6017038, 52646317, 21906754, 52644296, 86588542, 87168559, 265017, 265018, 265019, 264448, 264288, 264369, 52644296, 8658842, 21906766, 21906767, 21906768, 21906765, 21906767, 2486262, 25641129, 33657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657102, 35657113, 355770002
3133		87379414 (6265, 6266) Novel Protein sim. GBank gil4507613jreflNP_003738.1lpTNKS - TANKYRASE		polymerase	22278994, 22278998, 264905, 265006, 265007, 87168559, 264760, 21906767, 18108374, 22279000, 22279002, 264563
3134		94649816 (6267, 6268) Novel Protein sim. GBank gij1729827 sp P54633 TALA_DICDI - FILOPODIN (TALIN HOMOLOG)			264595, 264369, 264685, 264628, 264566
3135	•	86389356 (6289, 6270) Novel Protein șim. GBank gij3093478 (AF012927) - fibrinogen-binding protein (Streptococcus equi)		struct	22278996, 264095, 29331826, 33657402, 18108348, 263974
3136	94845839 (6271, 6272)	3136 94845839 (6271, 6272) Novel Protein sim. GBank gil627101 pirij544092 - probable carrier protein c2 - Caenorhabditis elegans	627101 pir  S44092 - probable   Contains protein domain (PF00153) - transport   Indication   Mitochondrial carrier proteins   Mitochondrial carrier   Mitochondria	transport	22278998, 264259, 264828, 265006, 265008, 60433438, 265019, 264764, 264288, 264769, 264689, 265020, 27486262, 263972, 65274791, 264557, 264558
3137		88257947 (6273, 6274) Novel Protein sim. GBank gij3342730 (AC005331) - R31341_1 [Homo sapiens]		UNCLASSIFIED	22276995, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 29331825, 29331826, 29331826, 29331826, 29331826, 29331827, 29331828, 265008, 21906754, 87168474, 265011, 87168559, 265017, 265018, 265019, 18108351, 264682, 264769, 21906765, 21906767, 21906769, 55811957, 35695917, 265020, 265021, 52844150, 18108374, 22279000, 22279002, 264482, 264488

### (#225503 (#277, #272)   Novel Protein sim: CBank gil/28928[m] 1814452C - Hyp-   (#C206858)   Novel Protein sim: CBank gil/28928[m] 1814452C - Hyp-   (#C206858]   Novel Protein sim: CBank gil/28928[m] 1814452C - Hyp-   (#52744)  (#6484 submit of signal recognition particle   (#52744)  (#6484 submit of signal recognition particle   (#527444)  (#6484 submit of signal recognition particle   (#527444)  (#6484 submit of signal recognition particle   (#52854)  Novel Protein sim: CBank gil/2498197 sp (39245) 5561   PiG     (#6285, #6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m     #64879         (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m         (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m         (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340287  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340387  m       (#6285, #6285, #6285, #6285)  Novel Protein sim: CBank gil/340387  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340387  m       (#6285, #6285, #6285)  Novel Protein sim: CBank gil/340387  m       (#6285, #6285, #6285, #6285)  Novel Protein sim: CBank gil/340387  m       (#6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285, #6285	2420		Mariet Brate: -:- Oberte :: 14 April 2014		
67225603 (6277, 6278) Novel Protein sim. GBank gij228938[pri]  1614452C - Hyp- rith glycoprotein [Zea diploperanis] 91222682 (6278, 6280) Novel Protein sim. GBank gij328938[pri]  1614452C - Hyp- (Z53744) 68tDA subuni of signal recognition particle (Z43744) 68tDA subuni of signal recognition particle (Z4323564 (6281, 6282) Novel Protein sim. GBank gij2289197[spi]056245[C561_PIG] (Z470CHROME B581 (CYTOCHROME B581 (CYTOCHROME B-561) (Z4743459) proteophosphosplospycan [Letshmania major] (Z4243459) proteophosphosplospycan [Letshmania major]			(AC006836) hypothetical protein [Arabidopsis thaliana]		33109954, 87168559, 264681, 264684,
6722562 (6277, 6278) Novel Protein sim. GBank gill28938[prill1814452C - Hyp- Inch gycoprotein [Zea dipoperatins] 91222682 (6278, 6280) Novel Protein sim. GBank gill32[amb[CAA37773] - Struct (Canis familiars] 67223564 (6281, 6283, Novel Protein sim. GBank gill3227 (AF035209) - putative (Canis familiars] 695419028 (6283, 6284) Novel Protein sim. GBank gill32037 (AF035209) - putative - CYTOCHROME B561 (CYTOCHROME B 561) - CYTOCHROME B561 (CYTOCHROME B 561)  (AL243459) proteophosphoglycan [Leishmania major] (AL243459) proteophosphoglycan [Leishmania major]					264685, 264686, 264687, 264768, 264688.
#7225503 (6277, 6278)   Novel Protein sim. CBank gij228938[pdf]   1814452C - Hyp-					264689, 264691, 264692, 264693, 33657109,
### 8722563 (#277, 6226) Novel Protein sin. GBank gij228938[pfl] 1814452C - Hyp-  ###################################					264631, 264634, 264635, 264636, 264637,
87323503 (8277, 6278) Novel Protein sim. GBank gilg28938[pril]1814452C - Hyp- ich glycoprotein [Zea diploperennis] 87223692 (6279, 6280) Novel Protein sim. GBank gilg321327 (AF035209) - pulative (X53744) 680A subunit of signal recognition particle (Zans familiaris) (X53744) 680A subunit of signal recognition particle (Zans familiaris)  87323564 (6281, 6282) Novel Protein sim. GBank gilg488197 [sp]098245[C561_Pri]  98419028 (6285, 6284) Novel Protein sim. GBank gilg4807 [sp]098245[C561_Pri]  - CYTOCHROME B561 (CYTOCHROME B 561)  (A.Z43459) proteophosphoglycan [Leistmania major]  (A.Z43459) Novel Protein sim. GBank gilg420387[emb](A846679.1] -					60170394, 83373044, 18108385, 18108388,
WOCASSIFED   Word Protein sim. GBank gil228938prill 181452C - Hyp-   WOCASSIFED   WORD group cate   Total sim. GBank gil228938prill 181452C - Hyp-   WOCASSIFED   Word Protein sim. GBank gil932lemb CAA3773] -	T				60432113, 22279000, 22279002
Inch givoprotein [Zea diploperennis]		87325503 (6277, 6278)	Novel Protein sim. GBank gij228938 prf  1814452C - Hyp-	UNCLASSIFIED	22278997, 22278998, 22278999, 264905,
91222692 (6279, 6280) Novel Protein sim. GBank gil932[amb]CAA37773] -  (X53144) 68tDA subunit of signal recognition particle [Caris familiaris]  87323564 (6281, 6282, 6282) Novel Protein sim. GBank gil3213227 (AF035209) - pulative			rich glycoprotein [Zea diploperennis]		265018, 265019, 21906765, 265020, 264636,
91222682 (6278, 6288 (6281, 6289, 6288) Novel Protein sim. GBank gil3213227 (AF035209) - pulative [Canis familiaris]  87323564 (6281, 6282, 6282) Novel Protein sim. GBank gil3213227 (AF035209) - pulative SNARE VI1 a [Mus musculus] SNARE VII	- 1	-			264557
(X53744) 68RbX subunit of signal recognition particle	_		Novel Protein sim. GBank gij932jemb CAA37773j -	struct	22278995, 56994075, 35696286, 264908,
(Canis familiaris     (Canis familiaris     (Canis familiaris     (Canis familiaris     V-SNARE VITTA   Mus musculus     V-SNARE VITTA   Mus musculus     85419028 (8283, 6284) Novel Protein sim. GBank gil243037   spiQ59245 C561_PIG     CYTOCHROME B561 (CYTOCHROME B-661)     CYTOCHROME B561 (CYTOCHROME B-661)     CYTOCHROME B561 (CYTOCHROME B-661)     CANOLASSIFIED     (AJ243459) proteophosphogiycan   Leishmania majori     CANOLASSIFIED     (AJ243459) proteophosphogiycan   Leishmania majori     CANOLASSIFIED				-	264909, 60433356, 21906754, 52644296,
87323564 (6281, 6282) Novel Protein sim. GBank gil3213227 (AF035209) - pulative UNCLASSIFIED v-SNARE VITIa [Mus musculus] 85419028 (6283, 6284) Novel Protein sim. GBank gil2498197 spj095245 C561_PIG cytochrome B561 (CYTOCHROME B-561) - CYTOCHROME B561 (CYTOCHROME B-561) 6A1243459) proteophosphogiycan [Leishmania major] UNCLASSIFIED (AL243459) proteophosphogiycan [Leishmania major]			[Canis familiaris]		87168474, 87168559, 264683, 264288,
87323564 (6281, 6282) Novel Protein sim. GBank gil3213227 (AF035209) - pulative  v-SNARE Vitta [Mus musculus]  96419028 (6283, 6284) Novel Protein sim. GBank gil5498197 spl095245 C561_PIG  - CYTOCHROME B561 (CYTOCHROME B-561)  65351475 (6285, 6285) Novel Protein sim. GBank gil5420387 emb[CAB46679.1] -  (A.1243459) proteophosphoglycan [Leistimania maiori]					264685, 264686, 265022, 264693, 27486262,
87323564 (6281, 6282) love Protein sim. GBank gij243227 (AF035209) - putative  v-SNARE Vita [Mus musculus]  95419026 (6283, 6264) Novel Protein sim. GBank gij2438197[sp]Q95245[C561_PIG  - CYTOC-HROME 8861 (CYTOCHROME B-561)  95351476 (6285, 6286) Novel Protein sim. GBank gij5420387[emb]CAB46679.1] -  (AJ243459) proteophosphogycan [Leistmania major]	1				35695855, 264630, 264555, 264566
V-SNARE Vitta [Mus musculus]		87323564 (6281, 6282)	Novel Protein sim. GBank gij3213227 (AF035209) - putative	UNCLASSIFIED	56182575, 35696286, 29331828, 264909,
95419028 (6283, 6284) Novel Protein sim. GBank gil2498197ispiQ95245[C561_PIG - CYTOCHROME B561 (CYTOCHROME B-561)  - CYTOCHROME B561 (CYTOCHROME B-561)  (AJ243459) protein sim. GBank gil5420387[emb]CAB46679.1] - (AJ243459) protein sim. GBank gil5420387[emb]CAB4679.1] - (AJ24346767[emb]CAB4679.1] - (AJ2434676767[emb]CAB4679.1] - (AJ24346767[emb]CAB4679.1] - (AJ24346767[emb]CAB4679.1] - (AJ24346767[emb]CAB4679.1] - (AJ2434767[emb]CAB4679.1] - (AJ2434767[emb]CAB4679.1			v-SNARE Vilta [Mus musculus]		265009, 265018, 18108351, 264369.
965419028 (6283, 6264) Novel Protein sim. GBank gil5420387 emb CAB46679.1   (AJZ43459) protein sim. GBank gil5420387 emb CAB46679.1   (AJZ43459) protein sim. GBank gil5420387 emb CAB46679.1 -					21906766, 29148627, 265020, 264628.
95419028 (8283, 6284), Novel Protein sim. GBank gij2498197lsp Q95245 C561_PIG - CYTOCHROME B561 (CYTOCHROME B-561) (AJ243459) Protein sim. GBank gij5420387 emb CAB46679.1  - UNCLASSIFIED  (AJ243459) proteophosphoglycan [Leistmania major]	L	_			264629, 264631, 18108385
- CYTOCHROME B561 (CYTOCHROME B-561)  95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 emb CAB46679.1  -  (AJ243459) proteophosphoglycan [Leishmania major]	~			cytochrome	52645156, 52646365, 22278995, 35696286,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420397 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]			- CYTOCHROME B561 (CYTOCHROME B-561)		22278998, 22278999, 60432049, 264259,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]					29331822, 29331824, 29331827, 29146499,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 jemb CAB45679.1  - (AJ243459) proteophosphogiycan [Leishmania major]					56182435, 265007, 60170831, 60432229,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 emb CAB45679.1 - (AJ243459) proteophosphoglycan [Leishmania major]					3355/402, 254595, 50433438, 254758,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387lemb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]					21906754, 264288, 264766, 264687,
95351475 (6285, 6286) Novel Protein sim. GBank gij5420387 emb CAB46679.1 - (AJ243459) proteophosphoglycan [Leishmania major]	_				52644229, 21906/65, 21906/67, 21906/68,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphogycan [Leisthmania major]					60170615, 52644150, 65274620, 33657109,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387lemb CAB46679.1  - (AJ243459) proteophosphogiycan [Leishmania major]					35695763, 18108370, 18108376, 65274791,
95351475 (6285, 6286) Novel Protein sim. GBank gij5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]		_			35695855, 264631, 264557, 87168518,
95351475 (6285, 6286) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]	1				60432113, 22279000
lycan [Leishmania major]				UNCLASSIFIED	264488, 56182575, 22278996, 22278998,
35596052, 29331828, 264508, 264905, 264907, 264907, 264907, 264901, 5264104, 56182404, 56182404, 56182404, 56182404, 56182404, 564901, 265018, 265908, 264901, 265018, 265908, 264901, 265908, 264901, 265908, 264901, 265908, 264901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901, 265901,			(AJ243459) proteophosphoglycan [Leishmania major]		22278999, 29331822, 29331824, 60432289,
264906, 264907, 264909, 52644044 56182435, 264511, 264512, 255008, 264904 60432222, 2555008, 264512, 2555008, 264514 60432222, 2655008, 264512, 2655008, 264760, 264685, 264684, 264788, 264685, 264684, 264788, 264885, 264885, 264885, 264885, 264885, 264885, 264885, 264885, 264885, 264885, 2648897, 264689, 35557023, 264693, 263957, 263957, 264693, 264897, 264638, 264897, 264639, 264897, 264639, 264639, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 26					35696052, 29331828, 264508, 264905,
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264486	. –				22279000, 22279002, 264563, 264566,
	-				264486

264488, 18108396, 22278996, 35696286, 22278997, 22278999, 29331826, 29331827, 35696052, 29331828, 264106, 265006, 265007, 265009, 33657402, 85588542, 265011, 18108351, 284448, 264369, 21906765, 21906766, 21906767, 26501, 265021, 265021, 35694150, 27486261, 18108370, 18108374, 35698423, 36182323, 83373044, 22279000, 22279002, 284567	18108397, 29331824, 29146499, 20281100, 265006, 55812038, 265010, 21906766, 29148627, 21906769, 29148784, 264692, 33657023, 33657109, 35695763, 263981, 56182323, 87168518	264259, 29331826, 29331828, 29331830, 264510, 264511, 265007, 265009, 264600, 265017, 18108351, 26448, 264369, 21906766, 265021, 264692, 33657109, 18108374, 35696423, 35695855, 60432113, 264564	56181686, 35696286, 60432049, 264259, 56182181, 29331825, 60432289, 35696052, 56182435, 265008, 264910, 60431735, 60433356, 60433438, 265010, 264448, 264288, 265022, 33657023, 33657109, 60431528, 65274791, 264631, 56182323, 264404, 22279002	35696286, 35696052, 264511, 85658542, 87168474, 264764, 35696423, 264555, 264556, 264557, 264558, 83373044, 56526486, 60432113	29331822, 35696052, 264109, 29148629, 18108381	264259, 29331822, 29331824, 29331825, 29331827, 52646317, 264686, 35695855, 56182323, 264639	29331822, 265008	21906754, 87168559, 264605, 21906768, 52644150, 27486264, 35696423, 22279000
	UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED	ерћ	transferase	опсодепе	UNCLASSIFIED
		Contains protein domain (PF00169) - PH domain	Contains protein domain (PF00702) - UNCLASSIFIED haloacid dehalogenase-like hydrolase		Contains protein domain (PF01363) - eph FYVE zinc finger	Contains protein domain (PF00043) - Itransferase Glutathione S-transferases.	Contains protein domain (PF00096) - oncogene Zinc finger, C2H2 type	
3144   95336329 (6287, 6288) Novel Protein sim. GBank gil4884468 emb CAB43322.1  - (AL050225) hypothetical protein [Homo sapiens]	86611657 (6289, 6290) Novel Protein sim. GBank gij3879709[emb]CAB03330] - (281118) Similarity to Human endosomal protein P162 (TR:Q15075); cDNA EST EMBL:Z14487 comes from this gene; cDNA EST EMBL:Z14556 comes from this gene; cDNA EST EMBL:D27011 comes from this gene; cDNA EST EMBL:D27015 comes from this gene; cDNA EST EMBL:D27015 comes from this gene; cDNA	87756314 (6291, 6292) Novel Protein sim. GBank gij2135746 pir  S69890 - mitogen   Contains protein domain (PF00169) - struct inducible gene mig-2 - human	94848512 (6293, 6294) Novel Protein sim. GBank gij3874279[emb]CAB07315.1] - (Z92825) predicted using Genefinder; cDNA EST yk315e12.3 comes from this gene; cDNA EST yk315e12.5 comes from this gene [Caenorhabditis elegans]	95362169 (6295, 6296) Novel Protein sim. GBank gi 5225322[gb AAD40851.1 AF08310 - (AF083108) sirtuin type 3 [Homo sapiens]	95308548 (6297, 6298) Novel Protein sim. GBank gil4200446 (AF102777) - FYVE finger-containing phosphoinoslitde kinase (Mus musculus)	Novel Protein sim. GBank gij3378454 jemb CAA76893  - (Y17850) gangiloside-induced differentiation associated protein 1 [Mus musculus]	87772355 (6301, 6302) Novel Protein sim. GBank gi[172591 (M63577) - SFP1 [Saccharomyces cerevisiae]	
95336329 (6287, 6288)	86611657 (6289, 6290)		94848512 (6293, 6294)	95362169 (6295, 6296)		87655472 (6299, 6300)		85698108 (6303, 6304)
4	3145	3146	3147	3148	3149	3150	3151	3152

			10000000000000000000000000000000000000		DEALES CORACRES RECEDENCE 20078006
3153	95317299 (6305, 6306)	3153   95317299 (6305, 6305) Novel Protein sim. CBank Januara (Protein domain (Protein Sim. CBank) Januara (Protein domain (Protein Sim. CBank) Januara (Protein domain (Protein Sim. CBank) Januara (CBank) J	WD domain G-beta repeat		22278997, 22278999, 60432049, 24£79999,
		4 Mis misculis		8	29331826, 60432289, 33656970, 264508,
				8	264908, 33657402, 264595, 60433438,
				82	87168474, 87168559, 264601, 265019,
				2	264448, 264682, 264764, 264288, 264369,
					264768, 21906765, 21906766, 21906767,
					21906768, 21906769, 29148784, 265021,
					265022, 60170615, 52644150, 264690,
				2	264691, 33657023, 65274620, 33657109,
			-		18108370, 35695855, 264638, 60170394,
				9	87168518, 60432113, 22279000, 22279002
3154	_	87718573 (6307, 6308) Novel Protein sim. GBank	4	ATPase_associated 2	22278998, 264259, 29331824, 66712502,
		qi 4680661 qb AAD27720.1 AF13294 - (AF132945) CGI-11			265008, 265010, 265017, 18108354, 264691.
		protein [Homo sapiens]			33657023, 264693, 20281149, 18108374
3155	87762394 (6309, 6310)	87762394 (6309, 6310) Novel Protein sim. GBank	-	UNCLASSIFIED	9331828, 264509, 264905, 264908, 264510,
		Gij728837 spjP39194jALU7_HUMAN - !!!! ALU SUBFAMILY		<u></u>	264511, 264512, 33657402, 264681, 264683,
		SQ WARNING ENTRY IIII			33657023, 18108370, 264634, 264639,
					18108385, 264563, 264486
3156		87737449 (6311, 6312) Novel Protein sim. GBank	Contains protein domain (PF00652) - Iransferase		56182575, 22278996, 22278997, 22278998,
		ail5630076lablAAD45821,11AC00601 - (AC006017) N-	Similarity to fectin domain of ricin		22278999, 60432049, 264259, 29331822,
		acetyloalactosaminyltransferase; similar to Q10473	beta-chain, 3 copies.		29331824, 66714117, 29331825, 29331826,
		(PID a1709559) [Homo sapiens]			29331827, 35696052, 52644045, 265007,
					265009, 60170831, 60432229, 60433356.
_					21906754, 33109954, 87168474, 265010.
					265017, 265018, 265019, 18108351, 264448,
					264288, 264689, 21906766, 21906768,
				<u></u>	21906769, 35695917, 265020, 265022,
	•				264692, 18108370, 35696423, 56182323.
					22279002
3157	88259577 (6313, 6314)				18108396, 264259, 29331826, 35696052,
					29146498, 87168559, 265017, 264448,
					264288, 264691, 18108366, 52645129,
	_				33080423, 32044332
3158		80034118 (6315, 6316) Novel Protein sim. GBank	Contains protein domain (PF00023) - kinase		264488, 263974
_		gi 5306064 gb AAD41895.1 AF15677 - (AF156778) ASB-3	Ank repeat		
3159	94124114 (6317, 6318)	3159   94124114 (6317   6318) Novel Protein sim. GBank gil5531272lemblCAB50897.11		UNCLASSIFIED	56182575, 22278999, 29331824, 264106,
}		[(AJ243800) WSC4 homologue [Kluvveromyces lactis]			60433356, 264758, 265011, 87168559,
					264448, 18108354, 264768, 21906768,
_					265020, 264691, 264692, 33657109,
					18108374, 35696423, 264555, 60170394.
					000001
3160		80221068 (6319, 6320) Novel Protein sim. GBank gij3930525 (AF064447) - sex-	Contains protein domain (PF00023) - struct		18108351, 264555, 264556, 264557, 264556, 264559
		מפופרתוות מנוסה ארסופות תסותים וס רפות נא וועום וווחסכיים א	Alla ichcol		

264488, 22278995, 22278997, 22278998, 264259, 29331822, 60432289, 29331828, 5264045, 265017, 265018, 264448, 264288, 21906764, 21906767, 265020, 18108374, 264636, 264566 3161 | 88074111 (6321, 6322)

Table 2

Tissue iD	Tissue Name	Tissue information	Disease Association
20281069	192xN	Protein-protein Interactions	Any
20281071	192xN	Protein-protein Interactions	Any
20281149	192xN	Protein-protein Interactions	Any
20281152	192xN	Protein-protein Interactions	Any
264111	276xN	Protein-protein Interactions	Any
264112	276xN	Protein-protein Interactions	Any
263966	384xN	Protein-protein Interactions	
263967	384xN	Protein-protein Interactions	Any
264110	552xN		Any
18108379	5PH 52.1 (Adrenal Gland)	Protein-protein Interactions	Any
10100379	SPR 32.1 (Adienal Gland)	Adrenai Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
18108381	5PH 52.2 (Fetal Lung)	Fetai Lung	Cystic Fibrosis, infection, lung cancer
18108383	5PH 52.3 (B's Lyphoma-Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108368	5PH 52.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108384	5PH 52.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders,
	,		neuropsychiatric disorders
18108394	SPH 53.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
18108355	5PH 53.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108359	5PH 53.3 (B's Lyphoma- Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108361	5PH 53.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108362	5PH 53.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108366	5PH 53.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders,
_	5111 55.0 (Etalli 111211105)		neuropsychiatric disorders
18108354	5PH 54.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia.
18108392	5PH 54.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108348	5PH 54.3 (B's Lyphoma- Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108382	5PH 54.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108395	5PH 54.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108365	5PH 54.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders,
	,		neuropsychiatric disorders
18108397	SPH 55.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
18108398	5PH 55.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108364	5PH 55.3 (B's Lyphoma- Raji)	Lymphoma derived from B cells	
18108388	SPH 55.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108358	5PH 55.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
20281099	5PH 56.2 (MG63)		
20281100	5PH 56.3 (UtSMC)		
264404	SPH.1 (Brain)	Whole Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia-telangiectasia, Leukodystrophies, Behavioral disorders,
		•	Addiction, Anxiety, Pain, Neuroprotection

264510	5PH.10 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264511	5PH.11 (Placenta)	Placenta	Infertility, birth defects
264512	5PH.12 (Thyroid)	Thyroid	Hyperparathyroidism, Hypoparathyroidism
264555	5PH.13 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
	(20110 11121011)	Done manow	thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host.
			management control of an area of the second
264556	5PH.14 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies.
			immunodeficiencies, transplantation, Graft vesus host,
	į		The state of the s
264557	5PH.15 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
	ì		thrombocytopenic purpura, autoimmume disease, allergies.
			immunodeficiencies, transplantation, Graft vesus host,
264558	5PH.16 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264559	5PH.17 (Bone Marrow)	Bone Marrow	Homobilio humano della disconti
204337	J. H. 17 (Bone Marrow)	Bolle Marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies.
			, , , , , , , , , , , , , , , , , , , ,
	İ		immunodeficiencies, transplantation, Graft vesus host,
264569	5PH.19 (One Fetal tissue and	Mixed	
201303	two cell lines)	"""	
264687	5PH.19.1 (fetal thymus -	Fetal Thymus	Hemophilia, hypercoagulation, Idiopathic
	CRL7046)		thrombocytopenic purpura, immunodeficiencies
264688	5PH.19.2 (hematopoetic stem	Hematopoeitic stem cells	Leukemia, osteoporosis, post-chemotherapeutic stem cell
	cells - CRL2043)		repopulation
264689	5PH.19.3 (osteogenic sarcoma cell lines - HTB96)	Osteogenic Sarcoma	Sarcomas, osteoporosis, osteopetrosis
264690	5PH.19.4 (Fetal Liver)	Fetal liver	Von Hippel-Lindau (VHL) syndrome.
			Cirrhosis, Transplantation
264691	5PH.19.5 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension,
			Congenital heart defects, Aortic stenosis, Atrial septal
			defect (ASD), Atrioventricular (A-V) canal defect, Ductus
			arteriosus, Pulmonary stenosis, Subaortic stenosis,
			Ventricular septal defect (VSD), valve diseases, Tuberous
			sclerosis, Scleroderma, Obesity, Transplantation
264602	SDI 10 6 (C-1)	6-1	10
264692	5PH.19.6 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
			thrombocytopenic purpura, Immunodeficiencies, Graft
264602	COVI 10 7 (Pinning)	District Control of the Control of t	vesus host
2 <b>64</b> 693	5PH.19.7 (Pituitary)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264482	SPH.2 (Brain)	Brain	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
		İ	-phopositive and syndrome, Muniple
			sclerosis Ataxia-
			sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders,

264600	5PH.21 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
		}	Epilepsy, Lesch-Nyhan syndrome, Multiple
	1		sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264601	5PH.22 (Bone Marrow)	Вопе Магтоw	Hemophilia, hypercoagulation, Idiopathic
		1	thrombocytopenic purpura, autoimmume disease, allergies,
1			immunodeficiencies, transplantation, Graft vesus host,
264602	5PH.23 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264603	5PH.24 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264604	5PH.25 (Lymph Node)	Lymph Node	Lymphedema, Allergies
264605	5PH.26 (Placenta)	Placenta	Infertility, birth defects
264634	5PH.28 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension,
			Congenital heart defects, Aortic stenosis, Atrial septal
	J		defect (ASD), Atrioventricular (A-V) canal defect, Ductus
	İ	Į.	arteriosus, Pulmonary stenosis, Subaortic stenosis,
			Ventricular septal defect (VSD), valve diseases, Tuberous
			sclerosis, Scleroderma, Obesity, Transplantation
264635	5PH.29 (Fetal Kidney)	Fetal Kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
		1	Interstitial nephritis, Glomerulonephritis, Polycystic
	- 1		kidney disease, Systemic lupus erythematosus, Renal
	1		tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
264483	5PH.3 (Bone Marrow)	Bone marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies,transplantation, Graft vesus host,
264636	5PH.30 (Lymph Node)	Lymph Node	Lymphedema , Allergies
264637	5PH.31 (P)ancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264638	5PH.32 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264639	5PH.33 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
		1	Parkinson's disease, Huntington's disease, Cerebral palsy,
		İ	Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
		1	telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264484	5PH.4 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264758	5PH.44.1 (Kidney)	Videou	Dishera Australian David
204/38	SETT.44.1 (MIGNEY)	Kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
			Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
364760	CDU 44 2 (Forel Liver)	Facilities.	Nyhan syndrome
264760	5PH.44.2 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome.
		L	Cirrhosis, Transplantation

264762	5PH.44.3 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension,
	,	1	Congenital heart defects, Aortic stenosis, Atrial septal
			defect (ASD), Atrioventricular (A-V) canal defect, Ductus
			arteriosus, Pulmonary stenosis, Subaortic stenosis,
	•	ļ	Ventricular septal defect (VSD), valve diseases, Tuberous
		İ	sclerosis, Scleroderma, Obesity, Transplantation
			scierosis, Scieroderma, Obesity, Fransplantation
264764	5PH.44.4 (Prostate)	Prostate	Prostate Cancer
264766	5PH.44.5 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
		1	thrombocytopenic purpura, Immunodeficiencies, Graft
			vesus host
264768	5PH.44.6 (pituitary)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
		ļ	disease, Stroke, Tuberous sclerosis, hypercalceimia,
	and the second		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
		<u> </u>	Addiction, Anxiety, Pain, Neuroprotection
264769	5PH.44.7 (Uterus)	Uterus	Infertility, birth defects
264905	5PH.48.1 (Burkitt's	Burkitt's Lymphoma	Lymphoma, blood cancers
264906	Lymphoma- Raji) 5PH.48.2 (Thalamus- Brain)	Thalamus	V- II - Li-t- (AII)
204700	JFM.46.2 (Thatamus- Brain)	i naiamus	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264007	CDVI 49 2 (Adams I Cl. 1)		
264907	5PH.48.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
264908	5PH.48.4 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
264909	SPH.48.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
264910	5PH.48.6 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
265006	SPH.50.1 (B's lymphoma)	Burkitt's Lymphoma	Lymphoma, blood cancers
265007	5PH.50.2 (thalamus)	Thalamus	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
	1 ' '		disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
		İ	Epilepsy, Lesch-Nyhan syndrome, Multiple
	-		sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
265008	5PH.50.3 (adrenal gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
265009	5PH.50.4 (fetal lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
265010	5PH.50.5 (salivary gland)	Salivary Gland	Dry mouth, infection
203010		Mammary Gland	Lactation disorders, breast cancer
265011	SPH.50.6 (mammary gland)	Immining Clair	
265011 18108385	5PH.51.1 (MCF-7)	Breast Cancer	Breast Cancer
265011 18108385 18108370	SPH.51.1 (MCF-7) SPH.51.2 (CCRF-CEM)	<del></del>	— <del>• • • • • • • • • • • • • • • • • • •</del>
265011 18108385 18108370 18108374	5PH.51.1 (MCF-7) 5PH.51.2 (CCRF-CEM) 5PH.51.3 (K-562)	Breast Cancer Cancer Cell line Cancer Cell line	Breast Cancer
265011 18108385 18108370 18108374 18108351	5PH.51.1 (MCF-7) 5PH.51.2 (CCRF-CEM) 5PH.51.3 (K-562) 5PH.51.4 (OVCAR-3)	Breast Cancer Cancer Cell line	Breast Cancer Cancer
265011 18108385 18108370 18108374 18108351 18108372	SPH.51.1 (MCF-7) SPH.51.2 (CCRF-CEM) SPH.51.3 (K-562) SPH.51.4 (OVCAR-3) SPH.51.5 (HL-60)	Breast Cancer Cancer Cell line Cancer Cell line Ovarian cancer Cancer Cell line	Breast Cancer Cancer Cancer Ovarian cancer Cancer
265011 18108385 18108370 18108374 18108351	5PH.51.1 (MCF-7) 5PH.51.2 (CCRF-CEM) 5PH.51.3 (K-562) 5PH.51.4 (OVCAR-3)	Breast Cancer Cancer Cell line Cancer Cell line Ovarian cancer	Breast Cancer Cancer Cancer Ovarian cancer Cancer Hemophilia, hypercoagulation, Idiopathic
265011 18108385 18108370 18108374 18108351 18108372	SPH.51.1 (MCF-7) SPH.51.2 (CCRF-CEM) SPH.51.3 (K-562) SPH.51.4 (OVCAR-3) SPH.51.5 (HL-60)	Breast Cancer Cancer Cell line Cancer Cell line Ovarian cancer Cancer Cell line	Breast Cancer Cancer Cancer Ovarian cancer Cancer

264508	5PH.8 (Fetal Brain)		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
	,		disease, Stroke, Tuberous sclerosis, hypercalceimia
			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Eniloper Local Multiple and desire Multiple
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia. Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264509 20798451	5PH.9 (Lymph Node)	Lymph Node	Lymphedema, Allergies
264487	5RH 56.3(UtSMC) 5RH.1 (Brain)		
204467	SKA. I (Brain)	Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
	1		Epilepsy, Lesch-Nyhan syndrome, Multiple
	1	1	sclerosis,Ataxia-
		į	telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264534	SRH.11 (Bone marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
	1		thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies,transplantation, Graft vesus host,
264535	5RH.12 (Bone marrow)	Воле Магтом	Hemophilia, hypercoagulation, Idiopathic
		1	thrombocytopenic purpura, autoimmume disease, allergies,
	İ		immunodeficiencies, transplantation, Graft vesus host,
			The state of the s
264563	5RH.19 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
	İ		Epilepsy, Lesch-Nyhan syndrome, Multiple
		1	sclerosis.Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264488	5RH.2 (Bone Marrow)	Вопе Магтом	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264564	5RH.20 (Lymph Node)	Lymph Node	Lymphedema, Allergies
264565	5RH.21 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264566	SRH.22 (Placenta)	Placenta	Infertility, birth defects
264567	5RH.23 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264591	5RH.25 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
		1	Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264592	5RH.26 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
		1	thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264593	5RH.27(thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264594	5RH.28 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264595	5RH.29 (Lymph Node)	Lymph Node	Lymphedema , Allergies
264489	5RH.3 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
	]	1	thrombocytopenic purpura, autoimmume disease, allergies,
	1	1	immunodeficiencies, transplantation, Graft vesus host,
			Inununodencies, transplantation. Graft vesus host

264596	5RH.30 (Placenta)	Placenta	Infertility, birth defects
264628	5RH.33 (fetal Kidney)	Fetal kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
			Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renai
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
264629	5RH.34 (lymph Node)	Lymph Node	Lymphedema, Allergies
264630	5RH.35 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264631	5RH.36 (thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264632	5RH.37 (Fetal Brain)	Fetal Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
		1	disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
ļ			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
		İ	Addiction, Anxiety, Pain, Neuroprotection
264490	SRH.4 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
	(2000 10222 00)		thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
			manufacturences, transplantation, Graft vesus nost,
264681	5RH.43.1 (fetal thymus -	Fetal Thymus	Hemophilia, hypercoagulation, Idiopathic
	CRL7046)		thrombocytopenic purpura, immunodeficiencies
264682	5RH.43.2 (hematopoetic stem	Hematopoeitic stem cells	Leukemia, osteoporosis, post-chemotherapeutic stem cell
	cells - CRL2043)		repopulation
264683	5RH.43.3 (osteogenic sarcoma	Osteogenic Sarcoma	Sarcomas, osteoporosis, osteopetrosis
·	cell lines - HTB96)		, , , , , , , , , , , , , , , , , , , ,
264684	5RH.43.4 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
			Cirrhosis, Transplantation
264685	5RH.43.6 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
			thrombocytopenic purpura, Immunodeficiencies, Graft
			vesus host
264686	SRH.43.7 (pituitary)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
	i		disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264757	5RH.44.1 (Kidney)	Kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
			Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
264759	5RH.44.2 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
			Cirrhosis, Transplantation
264761	5RH.44.3 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension,
			Congenital heart defects, Aortic stenosis, Atrial septal
			defect (ASD), Atrioventricular (A-V) canal defect, Ductus
			arteriosus, Pulmonary stenosis, Subaortic stenosis,
			Ventricular septal defect (VSD), valve diseases, Tuberous
			sclerosis, Scleroderma, Obesity, Transplantation
264763	5RH.44.4 (Prostate)	Prostate	Prostate Cancer
264765	5RH.44.5 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
	1		thrombocytopenic purpura, Immunodeficiencies, Graft
			vesus host

264767	5RH.44.6 (Pituitary)	Pituitary	No- transfer out to the second
		r nuna y	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
	1		disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy,
l			
	İ		Epilepsy,Lesch-Nyhan syndrome, Multiple sclerosis.Ataxia-
	1	1	telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264828	5DU 46 1 (1 b )/ 1 2	ļ	
264887	5RH.46.1 (Lymph Node)	Lymph Node	Lymphedema, Allergies
204887	5RH.47.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
18108377	SPU SO L (P's house)		Cirrhosis, Transplantation
18108380	5RH.50.1 (B's lymphoma)	Burkitt's Lymphoma	Lymphoma, blood cancers
10100300	5RH.50.2 (thalamus)	Thalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
İ			disease, Stroke, Tuberous sclerosis, hypercalceimia,
ł			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
1			telangiectasia, Leukodystrophies, Behavioral disorders,
L			Addiction, Anxiety, Pain, Neuroprotection
18108396	5RH.50.3 (adrenal gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,
18108391	5RH.50.4 (fetal lung)	Fetal Lung	Airway diseases, infection
18108357	5RH.50.5 (salivary gland)	Salivary Gland	Dry mouth, infection
18108390	5RH.50.6 (mammary gland)	Mammary Gland	Lactation disorders, breast cancer
264532	5RH.9 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
1			thrombocytopenic purpura, autoimmume disease, allergies,
Ì			immunodeficiencies,transplantation, Graft vesus host,
263974	736xN		
263976	736xN		
263981	736xN		
20281166	96xN		
20281169	96xN		
20281171	96xN		
263994	cDNA-ORF Selection		
264080	Mx96		
21906754	NQH 6.1 (HH729)		
22278996	NQH 6.10 (PrEC)	Endothelial cells	heart disease, cancer
22278997	NQH 6.11 (CAEC)	Endothelial cells	heart disease, cancer
22278998	NQH 6.12 (CSC)	Cancer Cell line	Cancer
22278999	NQH 6.13 (NHNPC)	Cancer Cell line	Cancer
22279000	NQH 6.14 (NHMC-RM)	Cancer Cell line	Cancer
22279002	NQH 6.15 (Hypothalmus)	Hypothalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
	1		telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection, Obesity
21906764	NQH 6.2 (In Dated Platelets)	Platelets	Clotting diseases, stroke
21906765	NOH 6.3 (HuVec)	Endothelial cells	heart disease, cancer
87168474	NQH 6.3 (Sized-HUVEC)	Endothelial cells	heart disease, cancer
21906766	NQH 6.4 (UtMVEC- myo)	Cancer Cell line	Cancer
21906767	NQH 6.5 (NHEM-neo)	Cancer Cell line	Cancer
21906768	NOH 6.6 (NHEK)	Cancer Cell line	Cancer
21906769	NOH 6.7 (ByCAEC)	Endothelial cells	heart disease, cancer
22278994	NQH 6.8 (NHA)	Cancer Cell line	Cancer
	1		Cancel

22278995	NQH 6.9 (PrSC)	Cancer Cell line	Cancer
27486261	NQH 7.1 (Jurkat E6-	Cancer Cell line	Cancer
	untreated)		
27486262	NQH 7.2 (TF1-untreated)	Cancer Cell line	Cancer
27486264	NQH 7.3 (U87-untreated)	Cancer Cell line	Cancer
27486265	NQH 7.4 (THP1-untreated)	Cancer Cell line	Cancer
29331822	NQH 8.1 (Brain- amygdala)		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
İ			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
Ì			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
ļ			telangiectasia, Leukodystrophies, Behavioral disorders,
i			Addiction, Anxiety, Pain, Neuroprotection
29331824	NQH 8.2 (Brain-		Von Hippel-Lindau (VHL) syndrome , Alzheimer's
	hippocampus)	1	disease, Stroke, Tuberous sclerosis, hypercalceimia,
	improve in the second		Parkinson's disease, Huntington's disease, Cerebral palsy,
İ			Epilepsy, Lesch-Nyhan syndrome, Multiple
!	1		sclerosis. Ataxia-
		}	telangiectasia, Leukodystrophies, Behavioral disorders,
l		1	Addiction, Anxiety, Pain, Neuroprotection
29331825	NOU 9 3 (Perio aubanasia		<u> </u>
29331825	NQH 8.3 (Brain- substantia		Von Hippel-Lindau (VHL) syndrome , Alzheimer's
	nigra)		disease, Stroke, Tuberous sclerosis, hypercalceimia,
]			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
ļ	1		telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
29331826	NQH 8.4 (small intestine)	Small intestine	digestive diseases, obesity, diabetes
29331827	NQH 8.5 (Spinal cord)	Spinal chord	paralysis, neurodegenerative disorders
29331828	NQH 8.6 (stomach)	Stomach	Stomach cancer
29331830	NQH 8.7 (Trachea)	Trachea	Airway diseases, infection
87168518	NQH 9.1 (Sized-MG-		
	63_treatment pool)		
87168559	NQH 9.2 (Sized-HEPG2		
	untreated)		
35695763	NQH.10.1 (MCF-7untreated)	Cancer Cell line	Cancer
35695855	NQH.10.2 (U-937_treatment	Cancer Cell line	Cancer
	pool)		
35695917	NQH.10.3 (JAR)	Cancer Cell line	Cancer
35696052	NQH.10.4 (PA-1)	Cancer Cell line	Cancer
35696286	NQH.10.5 (CADMEC)	Endothelial cells	heart disease, cancer
35696423	NQH.10.6 (CADMEC_LA)	Endothelial cells	heart disease, cancer
52644045	NQH.11.1 (SK-PN-DW)	Cancer Cell line	Cancer
52644150	NQH.11.2 (Chorionic Villus	Chorionic villus	fertility, birth defects
	Cells)		
52644229	NQH.11.3 (A549)	Cancer Cell line	Cancer
52644296	NQH.11.4 (U266B1)	Cancer Cell line	Cancer
52644332	NQH.11.5 (Daoy)	Cancer Cell line	Cancer
52644507	NQH.11.6 (SW1783)	Cancer Cell line	Cancer
52645080	NQH.12.1 (U-118MG)	Cancer Cell line	Cancer
52645129	NQH.12.2 (A204)	Cancer Cell line	Cancer
52645156	NQH.12.3 (T24)	Cancer Cell line	Cancer
52646317	NQH.12.4 (G-401)	Cancer Cell line	Cancer
52646365	NOH.12.5 (CaSki)	Cancer Cell line	Cancer
52646842	NOH.12.6 (SHP-77)	Cancer Cell line	Cancer
	Independent (Ages )	Compet cen une	l-centred

60424179	NQH.14.1 (Yale75_breast	Breast carcinoma	Breast Cancer
	carcinoma)	1	
60424269	NQH.14.2	Ovary tumor	Ovarian cancer
L	(Yale78B_ovarytumor)		
60431528	NQH.14.3	Prostate	Prostate Cancer
	(Yale79_prostateBPH)		
60431602	NQH.14.4	Prostate	Prostate Cancer
]	(Yale80_ProstateAdenocarcin		
	oma)		
60431735	NQH.14.5	Uterine Myoma	Uterine Cancer
50401050	(Yale86_UterineMyoma)		
60431850	NQH.14.6	Myometrium	Fertility
60432049	(Yale207_Myometrium) NQH.15.1 (Yale99_cervix)		
60432113		Cervix	Osteoporosis, cervical cancer
00432113	NQH.15.2 (Yale45 spleeniTP)		Hemophilia, Hypercoagulation, Idiopathic
	(Tate+3_spicentTF)		thrombocytopenic purpura, Immunodeficiencies, Graft vesus host
60432229	NQH.15.3 (Yale16 Skin)	Skin	
60432289	NQH.15.4 (Yale137_Parotid)	SKIR	wound healing, melanoma
32207	[1741112.4 (1 dieta /_ratotid)		
60433356	NOH.15.5	Small intestine	digestive diseases, obesity, diabetes
	(Yale38 SmallIntestine)	Jan Mestine	digestive diseases, obesity, diabetes
60433438	NOH.15.6	Colon	Colon cancer
	(Yale28_ColonAscending)		Colon Canada
65274444	NQH.17.1 (Larynx)	Larynx	Cancer
65274572	NQH.17.2 (Duodenum)	Duodenum	
65274620	NQH.17.3 (Kidney, Primary		Diabetes, Autoimmune disease, Renal artery stenosis,
	tumors)		Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
	1		tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
65274727	NQH.17.4 (Lung Pleura,	Lung	Airway diseases, infection
	normal)		
65274791	NQH.17.5 (Lung, Normal	Lung	Airway diseases, infection
2222244	Adult)		
83373044	NQH.18.230 (Pooled adrenal	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,
06660540	gland, placenta)	•••	
85658542	NQH.18.560 (Pooled uterus, BeWo pool)	Uterus	Infertility, birth defects
33656970	NQH.9.1 (MG-63 treatment	Conser Call Vise	
22020370	pool)	Cancer Cell line	Cancer
33657023	NQH.9.2 (HEPG2 untreated)		Von Hippel-Lindau (VHL) syndrome,
03037023	( Carrier of the carrier)		Cirrhosis, Transplantation
33657084	NQH.9.3 (PC3 untreated)	Cancer Cell line	Cancer
33657109	NQH.9.4 (TF-1 TPA)	Cancer Cell line	Cancer
33657182	NQH.9.5 (TF-1_TPO)	Cancer Cell line	Cancer
33657349	NQH.9.6 (TF-1_Hemin)	Cancer Cell line	Cancer
33657402	NQH.9.7 (HFDPC)	Cancer Cell line	Cancer
264259	NQH1(Mixture of eight adult		
	& two fetal tissues)		
264288	NQH2 (Ten tissues plus		
	lymphocyte control)		
264448	NQH3 (Bone Marrow)	Вопе Магтом	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
		•	immunodeficiencies, transplantation, Graft vesus host,
266015	1000		
265017	NQH4.1 (lymph node)	Lymph Node	Lymphedema, Allergies

265018	NQH4.2 (fetal kidney)	Fetal Kidney	IDiobase Australia D
205010	request (recall knamey)	retai Kidiley	Diabetes, Autoimmune disease, Renal artery stenosis,
	}	1	Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
		]	tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch
66712502	NQH4.2 (Sized)		Nyhan syndrome
265019	NQH4.3 (pituitary gland)	<del></del>	V. II. II. A. II.
203017	(pitutiary giaity)	1	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection, Obesity
66714117	NQH4.3 (Sized)	<del>-</del>	The state of the s
265020	NQH4.4 (testis)	testis	Infertility, birth defects
265021	NQH4.5 (fetal liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
	,	3.73.	Cirrhosis, Transplantation
265022	NQH4.6 (thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
18108376	NQH5.1 (MCF-7)	Breast cancer	Breast Cancer
18108387	NQH5.2 (CCRF-CEM)	Cancer Cell line	Cancer
264952	NRL1: HPLC		
	FRACTIONATION OF RE-		
	LIG		
263971	Old BB3 Baits		
263969	Old BB5 Baits		
263975	ORFSEL		
263972	OTHER Baits		
263978	pGALORF		
264106	PPBAITS		
264088	QC-YA7		
264089	QC-YA8		
264102	Resequenced Interactors		
264369	RRH.I		
60170394	RRH.10.1 (MCF-7untreated)	Breast cancer	Breast Cancer
50170615	RRH.10.2 (U-937_treatment	Cancer Cell line	Cancer
(0170031	pool)		
50170831 50174639	RRH.10.3 (JAR) RRH.11.8 (HeLa)	Cancer Cell line	Cancer
264113	rrOEA Baits	Cancer Cell line	Cancer
263973	RROEA B5 baits		
29146498	SRD 3.1 (SKMC)	Cancer Cell line	
29146499	SRD 3.2 (SKMC)	Cancer Cell line	Cancer
29147620	SRD 3.3 (RPTEC)	Cancer Cell line	Cancer
29147627	SRD 3.4 (HRCE)	Cancer Cell line	Cancer
29148629	SRD 3.6 (HRE)	Cancer Cell line	Cancer
29148784	SRD 3.7 (HRE)	Cancer Cell line	Cancer
5810764	SRD.7.1 (Lymph Node)	Lymph Node	Cancer
55811150	SRD.7.2 (pancreas)	Pancreas	Lymphedema, Allergies  Pancreatitis, diabetes, pancreatic cancer
5811386	SRD.7.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	1
	The state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the state of the s		Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,

55811576	SRD.7.4 (Pituitary Gland)	Diagram	
33811370	Sico. 7.4 (Filaliary Clarid)	Pituitary	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
		1	Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
İ			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection, Obesity
55811957	SRD.7.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
			Cirrhosis, Transplantation
55812038	SRD.7.6 (Fetal Kidney)	Fetal kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
			Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
56181562	SRD.8.1 (Lymph Node)	Lymph Node	Lymphedema, Allergies
56181686	SRD.8.2 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
56182181	SRD.8.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,
56182323	SRD.8.4 (Pituitary Gland)	Pituitary	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
		i	disease, Stroke, Tuberous sclerosis, hypercalceimia,
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
	1		Epilepsy, Lesch-Nyhan syndrome, Multiple
	1		sclerosis, Ataxia-
		ļ	telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection, Obesity
56182435	SRD.8.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome.
			Cirrhosis, Transplantation
56182575	SRD.8.6 (Fetal Kidney)		Diabetes, Autoimmune disease, Renal artery stenosis,
	1		Interstitial nephritis, Glomerulonephritis, Polycystic
			kidney disease, Systemic lupus erythematosus, Renal
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
	İ		Nyhan syndrome
32833986	SRD4: HL adapter		
56526486	SRD5.1:rr fragments		
33109954	SRD5: long-RXRJ		
5 <b>69</b> 94075	SRD9.1 (CS/SC)	Cancer Cell line	Cancer
263977	TSC Screen 1		

## Disclosed Sequences of ORFX Nucleic Acid and Polypeptide Sequences

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                             40
Val Leu Trp Pro Tyr Leu Leu Gln Phe Leu Thr Pro Val Arg Phe Thr
                                             60
Gly Ala Leu Thr Pro Leu Cys Arg Ser Leu Val His Leu Ala Gln Lys
                                         75
65
Arg Gln Glu Ala Gly Ala Asp Ala Phe Leu Ile Gln Tyr Asp Ala His
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Ala Ser Leu Pro Ser Pro Tyr Ala Val Thr Gly Arg Leu Leu Val Val
                                105
Ser Ser Ser Pro Tyr Leu Gly Asp Gly Arg Gly Ala Ala Ala Leu Arg
                            120
Leu Leu Ser Val Leu His Pro Asn Ile His Pro Leu Leu Gly Gln His
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Trp Glu Thr Thr Val Pro Leu Leu Leu Gly Tyr Leu Asp Glu His Thr
                                        155
                    150
Glu Glu Thr Leu Pro Gln Glu Glu Trp Glu Glu Lys Leu Leu Met Val
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Arg Ala Gly Val Arg Pro Ile Leu Gly Leu Lys Val Leu Ser Gly Leu
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Gly Gly Ala Gly Val Ala Glu Ala Gly Pro Pro Ala Ser Thr Ser Pro
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<213> Homo sapiens
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Cys Lys Pro Ala Val Gly Ser Ile Leu Ala Ser Cys Trp Asn Gln Pro
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Ile Met Asp Pro Ala Leu Val Pro Leu Gln Asp Thr Asn Asp Thr Phe
Met Ala Asn Met Gln Lys Asn Gly Thr Tyr Ser Ile Ile Pro Arg Ile
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Ala Gly Gly Glu Ile Thr Pro Asp Lys Leu Ile Ala Leu Gly Ala Val
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75
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                    70
Ala Lys Lys Tyr Asp Leu Tyr Thr Lys Ile Thr Gly Gly Gln Arg Ile
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Asp Leu Phe Gly Ala Gln Leu His Glu Leu Pro Gln Ile Trp Gly Glu
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Leu Val Asp Ala Gly Phe Glu Thr Gly
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acccaggtta gaatggtaaa ttgaaaggtg aatataaagg gagaatggtg aaatgaattt
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Arg Thr Lys Ala Leu Val Phe Phe Arg Ser Ser Thr Gly Asp Ser Asp
                            40
Ser Thr Ala Arg Ile Lys Lys Leu Ile Asn Gly Asn Ser Met Pro Val
Ala Glu Glu Leu Pro Trp Glu Met Ser His Thr Glu His Gln Ser Ser
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70

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Phe Pro Thr Pro Glu Ile Pro His Ser Leu Ala Pro Gly Thr Val Ala
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Val Cys Phe Ser Ser Ala Val Gln Ser Trp Ala Ile Arg Asn Thr Gly
                            40
Pro Leu Asn Thr Ser Pro Leu Leu Ala Leu Leu Leu Trp Ser Met
                        55
Gly Gly Val Gly Gly Ser Pro Val Ser Ala Pro Ala Ala Gly Ala His
                    70
Thr Thr Leu Ile Phe Gln Phe Trp Leu Trp Glu Pro Leu Pro Gln Val
                                   90
Ser Val Pro Gln Ala Pro Gly Leu Ser Phe Phe Tyr Cys Lys Ser Trp
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Lys Ser
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Ser Ala Met Tyr Thr Ser Val Glu Gly Trp Asp Tyr Val Asp Ser Leu
Tyr Phe Cys Phe Val Thr Phe Ser Thr Ile Gly Phe Gly Asp Leu Val
Ser Ser Gln His Ala Ala Tyr Arg Asn Gln Gly Leu Tyr Arg Leu Gly
Asn Phe Leu Phe Ile Leu Leu Gly Val Cys Cys Ile Tyr Ser Leu Phe
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Asn Val Ile Ser Ile Leu Ile Lys Gln Val Leu Asn Trp Met Leu Arg
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Lys Leu Ser Cys Arg Cys Cys Ala Arg Cys Cys Pro Ala Pro Gly Ala
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                            120
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<212> DNA
<213> Homo sapiens
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Gln Ser Phe Glu Glu Gly Ser Ser Gln Leu Cys Ile Phe Glu Gly Ser
Val Leu Leu Gly Pro Val Arg Ser Pro Val Gly Ser Arg Leu Glu
Trp Val Glu Pro Gly Gln Val Gly Ala Glu Thr Leu Glu Glu Ser Trp
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Ser Asn Ala Leu Cys Ile Trp Arg Gln Asn Gln Gln Arg Val Leu Trp
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Val Phe Ser Leu Thr Leu Ala Gly Trp Thr Val Val Pro Leu Asn
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gctgtttata cattaatgcc aatggttatg gctgatcaac acaggtctgt ttctgaacta
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<211> 111
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<213> Homo sapiens
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Glu Leu Pro Glu Asp Asn Glu Thr Ala Val Tyr Thr Leu Met Pro Met
                            40
Val Met Ala Asp Gln His Arg Ser Val Ser Glu Leu Leu Ser Asn Ser
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Lys Phe Asp Val Asn Tyr Ala Phe Gly Arg Val Lys Arg Ser Leu Leu
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His Ile Ala Ala Asn Cys Gly Ser Val Glu Cys Leu Val Leu Leu Leu
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                                    90
Lys Lys Gly Ala Asn Pro Asn Tyr Gln Asp Ile Ser Gly Cys Thr
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<213> Homo sapiens
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Lys Glu Ile Arg Gln Leu Pro Arg Tyr Asn Ile Thr Ile Lys Arg Val
Val Asn Met Thr Gly Lys Gly Arg Thr Pro Ser Trp Tyr Ser Leu Val
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Val Ala Gly Asn Gly Arg Gly Leu Val Gly Tyr Gly Glu Gly Lys Asp
Thr Asn Ile Ser Arg Ala Asn Lys Lys Ala Phe His Ala Ala Val Lys
Asn Met Asp Leu Val Ser Val His Arg Ser Lys Ser Gly Ala Asn Thr
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<213> Homo sapiens
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Gly Gly Ser Ala Ala Arg Phe Val His Ser Ala Gly Pro Thr Gly Ala
                            40
Arg Asn Arg Gln Glu Pro Pro Phe Pro Phe Glu Leu Ala Gly Arg Glu
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Pro Cys Thr Pro Arg Arg Ser Cys Ser Arg Pro Ala Ala Ala Leu Asp
                                        75
                    70
Pro Gly Ile Ser Ala Leu Ser Gly Ala Gln Glu Ala Ser Leu Thr Arg
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Arg Leu Val Ser Ala Cys Ser Arg Ser Ser Pro Leu Leu Ala Pro Thr
                                105
Ser Ile Ser Glu Gln Ser
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240
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Leu Trp Ala Thr Ile Glu Gln Asp Leu Leu Thr Lys Gly Asp Glu Cys
Lys Phe Gly Gly lys Ser Val Arg Asp Gly Met Ala Gln Ser Gly
                            40
Thr Ala Thr Arg Asp Asn Pro Asn Val Leu Asp Phe Val Ile Thr Asn
Val Met Ile Ile Asp Ala Lys Leu Gly Ile Ile Lys Ala Asp Ile Gly
Ile Arg Asp Gly Arg Ile Val Gly Ile Gly Gln Ala Gly Asn Pro Asp
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Thr Met Asp Asp Val Thr Pro Asn Met Ile Ile Gly Ala Ser Thr Glu
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                            105
Val His Asn Gly Ala
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<211> 355
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355
<210> 36
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Pro Trp Gln Ala Gly Ala Trp Leu Val Pro Gly Ser Pro Ala Cys Thr
                            40
Ser Val Arg Pro Pro Asp Lys Ser Ser Pro Pro Pro Gly Thr Pro Gly
Pro Thr Gly Trp Gly Leu Leu Pro Gln Arg Val Asp Thr Gly Arg Arg
                    70
                                        75
Glu Cys Ser Ala His Ser Pro Pro Leu Ala Gln Ala Pro Trp Leu Gly
                                    90
Ala Gly Pro Arg Pro Thr Leu Gly Arg Ala Gly Gly Ala Gly Arg Ala
                                105
            100
Thr Ala Ser Leu His Ala
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492
<210> 38
<211> 127
<212> PRT
<213> Homo sapiens
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Ala Gly Tyr Arg Pro Arg Val Leu Ala Asp Asp Val Cys Cys Gly Leu
Thr Trp Ile Thr Thr Gly Gln Leu Asp Gly Ala Arg Arg Arg Leu Arg
                            40
Ala Gly Leu Asp Val Leu Ala Pro Leu Ser Asp Ala Ser Val Pro Val
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                        55
                                             60
Val Gly Leu Glu Pro Ser Cys Thr Thr Val Trp Arg Asp Asp Ala Leu
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Arg Leu Leu Pro Asp Asp Pro Arg Val His Arg Val Ala Arg Asn Met
His Thr Val Ala Glu Met Leu Glu Ala Ala Gln Trp Thr Pro Pro Ser
                                105
Leu Ala Gly His Thr Leu Val Ala Gln Pro His Cys His Pro Ala
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<210> 39
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<213> Homo sapiens
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120
gaaatggaag acgtttatta cagcattgcc ggaaaacaac tggtgagcaa cttctctgcg
caagtcatgc gtggtgataa aattgcgctg attggcccga acggttgtgg taaaacgacg
ttgctgaaac tgatgttaag taagattcag gcagacagcg gccgtgttca ctgcggtact
aaactggaag ttgcgtactt cgaccagcac cgtgctgagc tggatcctga gcgtacggtg
atggataacc tggccgaagg taagcaggaa gtgatggtaa atggccgtgt an
412
<210> 40
<211> 137
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<213> Homo sapiens
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Asp Arg Arg Glu Val Met Xaa Thr Ala Lys Met Gln Val Val Glu Ala
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Ala Ser Ser Gly Lys Ile Val Phe Glu Met Glu Asp Val Tyr Tyr Ser
Ile Ala Gly Lys Gln Leu Val Ser Asn Phe Ser Ala Gln Val Met Arg
Gly Asp Lys Ile Ala Leu Ile Gly Pro Asn Gly Cys Gly Lys Thr Thr
                    70
Leu Leu Lys Leu Met Leu Ser Lys Ile Gln Ala Asp Ser Gly Arg Val
                                    90
His Cys Gly Thr Lys Leu Glu Val Ala Tyr Phe Asp Gln His Arg Ala
                                105
Glu Leu Asp Pro Glu Arg Thr Val Met Asp Asn Leu Ala Glu Gly Lys
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Gln Glu Val Met Val Asn Gly Arg Val
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135

130

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Glu Arg Thr Val Ala Lys Asp Phe Val Thr Thr Glu Val Glu Pro Met
                            40
Trp Asp Ala Ala Asp Val Met Arg Met Gly Lys Asp Leu Phe Ile Gln
                        55
His Gly Leu Thr Thr Asn Arg Lys Ser Met Glu Trp Phe Lys Arg Tyr
Tyr Pro Asp Phe Arg Val His Ala Val Asn Phe Pro Gly Asp Pro Tyr
                                    90
                85
Pro Ile His Ile Asp Ala Thr Phe Val Pro Leu Arg Pro Gly Leu Ile
                                105
Ile Asn Asn Pro Asn Arg Pro Leu Pro Gln Glu Gln Arg Lys Ile Phe
                            120
Glu Ala Asn Asp Trp Gln Ile Val Asp Ala Ala Gln Pro Ala His Asp
                        135
                                            140
Thr Pro Pro Glu Leu Cys Tyr Ser Ser Val Trp Leu Ser Met Asn Cys
                    150
                                        155
Leu Val Leu Asp Pro Lys Thr Val Ile Cys Glu Ala Ser Glu Val His
                                    170
Gln Met Glu Gln Met Asp Lys Leu Gly Met Asn Val Ile Pro Val Ala
                                185
            180
Phe Arg Asp Ala Tyr Pro Phe Gly Gly Gly Leu His Cys Ala Thr Ala
                            200
Asp Val Tyr Arg Glu Gly Thr Cys Glu Asp Tyr Phe Pro Asn Gln Val
                        215
                                            220
Asp Asp Pro Thr Leu Val
225
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<212> DNA
<213> Homo sapiens
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gtgaccacct ggtggggaat aaggcgcttc tgggacatag aggctgcctt ccagctgcgc
ctggcagagc tgttgacaca acagcatggt ctgcagtgcc gggccactgc cacgcacacc
gatgtccttt aaggatggat ttgggttttc ggattcgcgt ggcctatcag cgggagtccc
agatectgaa ggaagtgeag ageceagagg ggatgatete getgagggae acagetgeet
coetcoqeet tgagagagac acaaggeagt tgccactget caccagtgee etgcacgn
358
<210> 44
<211> 105
<212> PRT
<213> Homo sapiens
<400> 44
Met Glu Cys Gln Glu Val Gly Asp His Leu Val Gly Asn Lys Ala Leu
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1
Leu Gly His Arg Gly Cys Leu Pro Ala Ala Pro Gly Arg Ala Val Asp
                                25
Thr Thr Ala Trp Ser Ala Val Pro Gly His Cys His Ala His Arg Cys
Pro Leu Arg Met Asp Leu Gly Phe Arg Ile Arg Val Ala Tyr Gln Arg
Glu Ser Gln Ile Leu Lys Glu Val Gln Ser Pro Glu Gly Met Ile Ser
                                        75
Leu Arg Asp Thr Ala Ala Ser Leu Arg Leu Glu Arg Asp Thr Arg Gln
                                    90
                85
Leu Pro Leu Leu Thr Ser Ala Leu His
            100
                                105
<210> 45
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<212> DNA
<213> Homo sapiens
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geggeteetg gaateceaga geagtatggt ggegaeggtg eggatgegat tgegteegea
ataatcatgg aagaggtcgc tcgagtctgt gcgtcgtcgt ccaccgtcat atcgtccaat
gagettggta eegteeetet eeteaaatae ggtagegagg ageagaggaa aegttatett
totgaagttg ottogggtaa ggoactttto ggatatgogo totocgaggo tgatgotgga
tcagatccag ctgcacttaa gtgtcgagcc gacgaagatg gggacagttt cgtcctgaat
360
ggcgttaagg cttgggtcac ggaggctggc gaggccaagt acctggtgat atttgcggtt
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qqcattagct acggggctcc ggagcacaaa atggggatac gcgggtcagt taccagggaa
gtggttttca agaatacgcg tatccccaag gaacgagtaa ttggccgtcg agggcacggt
600
ctgagtgttg ctctaggtac gcttgataac tctcgtgtct cgattgctgc tcaagcagtg
ggaattgccc aaggagcttt agacattgcc acggattacg tccagaagcg caagcagttt
ggccagccac tgtccaattt tgagggaatc cagttcatgc tcgcagacat ggcaatgcgt
780
ttggaggcgg cgcgagcgct gacatactct gcagctgatc gtagtgggcg ccagactgac
gatgtgagtt actteggege ggeggeeaaa tgtttegett eegacacage gatggeagtg
900
tgcac
905
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<210> 46

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<211> 301
<212> PRT
<213> Homo sapiens
<400> 46
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Glu Ala Gly Phe Ala Ala Pro Gly Ile Pro Glu Gln Tyr Gly Gly Asp
Gly Ala Asp Ala Ile Ala Ser Ala Ile Ile Met Glu Glu Val Ala Arg
                            40
Val Cys Ala Ser Ser Ser Thr Val Ile Ser Ser Asn Glu Leu Gly Thr
                        55
Val Pro Leu Lys Tyr Gly Ser Glu Glu Gln Arg Lys Arg Tyr Leu
Ser Glu Val Ala Ser Gly Lys Ala Leu Phe Gly Tyr Ala Leu Ser Glu
Ala Asp Ala Gly Ser Asp Pro Ala Ala Leu Lys Cys Arg Ala Asp Glu
                                105
Asp Gly Asp Ser Phe Val Leu Asn Gly Val Lys Ala Trp Val Thr Glu
                            120
Ala Gly Glu Ala Lys Tyr Leu Val Ile Phe Ala Val Thr Asp Pro Asp
                        135
Asp Pro Arg His Arg Ile Ser Ala Leu Met Val His Ala Asp Asp Pro
                   150
                                        155
Gly Ile Ser Tyr Gly Ala Pro Glu His Lys Met Gly Ile Arg Gly Ser
                                    170
               165
Val Thr Arg Glu Val Val Phe Lys Asn Thr Arg Ile Pro Lys Glu Arg
                               185
Val Ile Gly Arg Arg Gly His Gly Leu Ser Val Ala Leu Gly Thr Leu
                            200
Asp Asn Ser Arg Val Ser Ile Ala Ala Gln Ala Val Gly Ile Ala Gln
                        215
Gly Ala Leu Asp Ile Ala Thr Asp Tyr Val Gln Lys Arg Lys Gln Phe
                   230
                                        235
Gly Gln Pro Leu Ser Asn Phe Glu Gly Ile Gln Phe Met Leu Ala Asp
                                   250
               245
Met Ala Met Arg Leu Glu Ala Ala Arg Ala Leu Thr Tyr Ser Ala Ala
                                265
Asp Arg Ser Gly Arg Gln Thr Asp Asp Val Ser Tyr Phe Gly Ala Ala
                            280
Ala Lys Cys Phe Ala Ser Asp Thr Ala Met Ala Val Cys
                        295
<210> 47
<211> 379
<212> DNA
<213> Homo sapiens
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atgeatetta eegetgegta tgeegtaget aeggaagetg ggtgeeatat eeggttaagt
120
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cagtatgete ggaaagteeg ecagaegeag ttaagagtgg aatacetgeg cetteggetg
gegageetge etggtggtga tgetggegeg geagtaggaa ttgategteg aetgegttta
240
gatttcgaaa aaggactcac caaatcccag ggtcgacgag aagagttcat acccgtcggc
300
gaagacgcca gcacgtataa cagacttatg aaagcgctgc gccaacgcca tgatgtcatc
aaatccggaa agcttgccc
379
<210> 48
<211> 106
<212> PRT
<213> Homo sapiens
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Met His Leu Thr Ala Ala Tyr Ala Val Ala Thr Glu Ala Gly Cys His
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Ile Arg Leu Ser Gln Tyr Ala Arg Lys Val Arg Gln Thr Gln Leu Arg
                                25
Val Glu Tyr Leu Arg Leu Arg Leu Ala Ser Leu Pro Gly Gly Asp Ala
                            40
Gly Ala Ala Val Gly Ile Asp Arg Arg Leu Arg Leu Asp Phe Glu Lys
Gly Leu Thr Lys Ser Gln Gly Arg Arg Glu Glu Phe Ile Pro Val Gly
                                        75
                    70
Glu Asp Ala Ser Thr Tyr Asn Arg Leu Met Lys Ala Leu Arg Gln Arg
                85
His Asp Val Ile Lys Ser Gly Lys Leu Ala
            100
<210> 49
<211> 309
<212> DNA
<213> Homo sapiens
<400> 49
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atccctctaa tttttgtgtc tccttctgta tcatcaaatt ttccctctct actgagtctc
120
ttgcatctcc ttggaagcat gctgtactat gtcccatcct taaagaactc cccttgtctg
cacattaccc tetgecaget ggeteatttt tetgeteece tttacaggga aactetteaa
aaagttatct ccacctcctt ccatctcatg ttctcttgaa cctgcagtac tgggtgctcc
300
ctccttttg
309
<210> 50
<211> 101
<212> PRT
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<213> Homo sapiens

<400> 50 Met Met Leu Ala Trp Thr Ile Leu Val Pro Val Pro Leu Ser Pro Ala 10 Glu Gly His Pro Ser Asn Phe Cys Val Ser Phe Cys Ile Ile Lys Phe 25 Ser Leu Ser Thr Glu Ser Leu Ala Ser Pro Trp Lys His Ala Val Leu Cys Pro Ile Leu Lys Glu Leu Pro Leu Ser Ala His Tyr Pro Leu Pro 55 Ala Gly Ser Phe Phe Cys Ser Pro Leu Gln Gly Asn Ser Ser Lys Ser 75 Tyr Leu His Leu Leu Pro Ser His Val Leu Leu Asn Leu Gln Tyr Trp Val Leu Pro Pro Phe 100 <210> 51 <211> 512 <212> DNA <213> Homo sapiens <400> 51 agatetttqa agaattqcca cactqtcttc ctccctgctt ataatttcct tattccctag gatgtgatec ttgttettgg ggeeteacat ggeagetgga tetetggega ttgeatetga gttccagaca ccaggatgga aaagaaaaga aggaggggca agaggaaccc ccagatgctc 180 cttaagagct actgcgtggc attcccactt gcatctcatt tgctcgatcg ctgtcactgt gccctaacga gctgcaagga cactggggaa atgagtctgt cttgtacttc atgtgcccct caaaatcttc tgttgctgag ggagaagagg ccagccggta ttgaggaaca actagcactt tetgetteeg egteecaggg ggaegtgggt gtgttgaate cacacegggg gtgeggaeet ctgaggctgg gctggatggg acatcaggtg ggccctctgt ttcatttatg tgacctccca tcaggtcttc tggttggatc ctgctttcta ga 512 <210> 52 <211> 125 <212> PRT <213> Homo sapiens <400> 52 Met Glu Lys Lys Arg Arg Arg Gly Lys Arg Asn Pro Gln Met Leu Leu 10 Lys Ser Tyr Cys Val Ala Phe Pro Leu Ala Ser His Leu Leu Asp Arg Cys His Cys Ala Leu Thr Ser Cys Lys Asp Thr Gly Glu Met Ser Leu

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35
Ser Cys Thr Ser Cys Ala Pro Gln Asn Leu Leu Leu Arg Glu Lys
                        55
                                            60
Arg Pro Ala Gly Ile Glu Glu Gln Leu Ala Leu Ser Ala Ser Ala Ser
                    70
                                        75
Gln Gly Asp Val Gly Val Leu Asn Pro His Arg Gly Cys Gly Pro Leu
Arg Leu Gly Trp Met Gly His Gln Val Gly Pro Leu Phe His Leu Cys
                                105
Asp Leu Pro Ser Gly Leu Leu Val Gly Ser Cys Phe Leu
                            120
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<212> DNA
<213> Homo sapiens
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aaaattcqat cccgtcgtgc gtcgtcacat tgagttcaag gaggcccgct aatggccaaa
aagtccaaga ttgtcgccca gaagaaacgt gagaagctcg tagcccaata cgccgaaagg
cgcgccgaac tcaaggccat catgaagtgc ccaactgcct cattggacga acgcatggag
geategegta agetgteteg cetgeegege gatteatece eegtgeggtt aegtaacegt
qaccaagteg acgggcgtec ccgcggctac gttggcaagg ccggtgtgtc ccgtatccgt
ttccgtgaga tggcccaccg cggcgaactc cccggaatcg cgaagtcaag ctggtgaagc
catggcagta ccgaagcgaa agaagtcccg ttcgaccacg cgtcataggc gggc
474
<210> 54
<211> 101
<212> PRT
<213> Homo sapiens
<400> 54
Met Ala Lys Lys Ser Lys Ile Val Ala Gln Lys Lys Arg Glu Lys Leu
Val Ala Gln Tyr Ala Glu Arg Arg Ala Glu Leu Lys Ala Ile Met Lys
Cys Pro Thr Ala Ser Leu Asp Glu Arg Met Glu Ala Ser Arg Lys Leu
Ser Arg Leu Pro Arg Asp Ser Ser Pro Val Arg Leu Arg Asn Arg Asp
Gln Val Asp Gly Arg Pro Arg Gly Tyr Val Gly Lys Ala Gly Val Ser
                    70
Arg Ile Arg Phe Arg Glu Met Ala His Arg Gly Glu Leu Pro Gly Ile
                                    90
Ala Lys Ser Ser Trp
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100

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<210> 55
 <211> 378
 <212> DNA
 <213> Homo sapiens
 <400> 55
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 teggegeage caageeegea gegtgetgee aggegeaage gacaaacace ggeeegtggg
 tggtgttcga ccatgtgcgt tgcacccacg acacctttct gatcgacgtc tttctcaacc
 agecegatge caeegegeag caggicaatg cegacaacce geactaegte gggegtttea
               4 10
 geogeategg catgggeetg gtggatgaca agggeegttg cattacccag ggegtatege
 gegegttgaa tgeggegege ageaceaagg egetgaacet gggaeegagt gaegeggege
 agttatcggt gaggcgta
 378
 <210> 56
' <211> 125
 <212> PRT
 <213> Homo sapiens
 <400> 56
 Met Ala Gln Asp Ser Arg His Ile Gly Tyr Asp Tyr Gly Thr Pro Val
 Ala Pro Gln Phe Gly Ala Ala Lys Pro Ala Ala Cys Cys Gln Ala Gln
                                 25
 Ala Thr Asn Thr Gly Pro Trp Val Val Phe Asp His Val Arg Cys Thr
                             40
 His Asp Thr Phe Leu Ile Asp Val Phe Leu Asn Gln Pro Asp Ala Thr
                         55
                                              60
 Ala Gln Gln Val Asn Ala Asp Asn Pro His Tyr Val Gly Arg Phe Ser
                                         75
                     70
 Arg Ile Gly Met Gly Leu Val Asp Asp Lys Gly Arg Cys Ile Thr Gln
                 85
                                     90
 Gly Val Ser Arg Ala Leu Asn Ala Ala Arg Ser Thr Lys Ala Leu Asn
             100
                                 105
 Leu Gly Pro Ser Asp Ala Ala Gln Leu Ser Val Arg Arg
                             120
 <210> 57
 <211> 388
 <212> DNA
 <213> Homo sapiens
 agacccaccc gacacagatc aggagtegtc atgtccagaa agaagaaggt cggcatcctc
 60
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accgcaggcg gtgattgccc cgggctcaac gccgctatcc gcggatttgg caaggctgcc
atccgccagc acgacatgga gctcatcggt attcaggacg gctttcttgg attggcggga
aaccgcacca tetecettgg eccgegtgee eteteaggea tettgaeggt eggegggaee
atcctgggaa ctagccgtga caaggtcaat cacatgatta tcgacggcga ggaacgggat
atggtcccca ccaccgtcga gaattacgag aagctggggc ttgacgcttt ggtgactttg
ggtggcggtg gcaccgccaa gaacgcgt
388
<210> 58
<211> 129
<212> PRT
<213> Homo sapiens
<400> 58
Arg Pro Thr Arg His Arg Ser Gly Val Val Met Ser Arg Lys Lys
Val Gly Ile Leu Thr Ala Gly Gly Asp Cys Pro Gly Leu Asn Ala Ala
Ile Arg Gly Phe Gly Lys Ala Ala Ile Arg Gln His Asp Met Glu Leu
Ile Gly Ile Gln Asp Gly Phe Leu Gly Leu Ala Gly Asn Arg Thr Ile
                        55
Ser Leu Gly Pro Arg Ala Leu Ser Gly Ile Leu Thr Val Gly Gly Thr
                                        75 .
                    70
Ile Leu Gly Thr Ser Arg Asp Lys Val Asn His Met Ile Ile Asp Gly
                                    90
Glu Glu Arg Asp Met Val Pro Thr Thr Val Glu Asn Tyr Glu Lys Leu
Gly Leu Asp Ala Leu Val Thr Leu Gly Gly Gly Gly Thr Ala Lys Asn
                            120
Ala
<210> 59
<211> 417
<212> DNA
<213> Homo sapiens
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ggtaccatcg gagetegaca agaaatggtt gggtgaagte gtggettetg etceacceag
60
tgccctcatg ggtcagccca cctgaatatc ttcatgcctg tgcatttctc ctgatgttca
cgtgtgccct gtgtttttac gcatctgtga tcgtgcaccc acgcgtctca gagaggagcc
180
cgtttgggaa tccggagaat gtgcgctggc ggaagagcgt cacacactgg aagcaaacct
cagaccgcgt ggacaagacc aaggatgaaa tggaacacga ggccttggtg gaagggaacc
300
```

```
tggcaaccga ggcaaqccta gtggttctgg acacactgga gatcatcgtg cagacggtga
tgctttcaqa aqcccqqqaq aqcqtcttqq qqgcaqtqct qaaqgttqtg ctgtaca
417
<210> 60
<211> 101
<212> PRT
<213> Homo sapiens
<400> 60
Met Phe Thr Cys Ala Leu Cys Phe Tyr Ala Ser Val Ile Val His Pro
                                     10
Arg Val Ser Glu Arg Ser Pro Phe Gly Asn Pro Glu Asn Val Arg Trp
                                25
Arg Lys Ser Val Thr His Trp Lys Gln Thr Ser Asp Arg Val Asp Lys
                            40
Thr Lys Asp Glu Met Glu His Glu Ala Leu Val Glu Gly Asn Leu Ala
Thr Glu Ala Ser Leu Val Val Leu Asp Thr Leu Glu Ile Ile Val Gln
                    70
                                         75
Thr Val Met Leu Ser Glu Ala Arg Glu Ser Val Leu Gly Ala Val Leu
                                    90
Lys Val Val Leu Tyr
            100
<210> 61
<211> 304
<212> DNA
<213> Homo sapiens
<400> 61
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gcacacatat ttgcaggctt ggagagagtg tgtgggggca tgtactttcg gtgggtcaag
tatgaagaag caggccttat aaacacatat tctgacctta acctgtactt cagaagagga
ccgctgactc accaaggagg cctgaaggac aaggcagcat ctctgtcttc acatgagtcc
toccotagae egggeceatg gecaggeetg accaeagage teccattgee ttteetgeae
300
gcgt
304
<210> 62
<211> 92
<212> PRT
<213> Homo sapiens
<400> 62
Met Gly Ala Leu Gln Phe Trp Arg Ser Leu Ser Ala His Ile Phe Ala
                 5
                                    10
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Gly Leu Glu Arg Val Cys Gly Gly Met Tyr Phe Arg Trp Val Lys Tyr
                    25
Glu Glu Ala Gly Leu Ile Asn Thr Tyr Ser Asp Leu Asn Leu Tyr Phe
Arg Arg Gly Pro Leu Thr His Gln Gly Gly Leu Lys Asp Lys Ala Ala
                                            60
                        55
Ser Leu Ser Ser His Glu Ser Ser Pro Arg Pro Gly Pro Trp Pro Gly
                    70
Leu Thr Thr Glu Leu Pro Leu Pro Phe Leu His Ala
<210> 63
<211> 577
<212> DNA
<213> Homo sapiens
<400> 63
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ctgacggtgc tagctgggtg gctcacccta gccgggcgta tcagtgtcgg ggaactcgtc
acceptage georgecca aaccetegge cetecgetge gageactagg egtegacace
gcgacgatgt tggccaccgc ccacgcctcc ggggaccgat tctgtgagtt gcgtgatagc
coggoaquet ggcagateca coccgaegac ggtgcccgca ccacaccggg tgatggcccg
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gtgctggtgg ctccccgctc ccccgaactg ttcgacgata ctgcccgtgc gaacatcgtg
cttgacagcc agacgactgt cgccaggctg aatgcat
577
<210> 64
<211> 192
<212> PRT
<213> Homo sapiens
<400> 64
Arg Val Lys Gly Val Tyr Thr Gly Thr Ile Asn Ala Ser Val Gly Val
Phe Ile Thr Ala Leu Thr Val Leu Ala Gly Trp Leu Thr Leu Ala Gly
Arg Ile Ser Val Gly Glu Leu Val Thr Val Val Gly Leu Ala Gln Thr
                            40
Leu Gly Pro Pro Leu Arg Ala Leu Gly Val Asp Thr Ala Thr Met Leu
                        55
Ala Thr Ala His Ala Ser Gly Asp Arg Phe Cys Glu Leu Arg Asp Ser
                                                             80
                    70
                                        75
```

```
Pro Ala Ala Trp Gln Ile His Pro Asp Asp Gly Ala Arg Thr Thr Pro
                                    90
                85
Gly Asp Gly Pro Val Glu Leu His Ile Pro Val Arg Asp Phe Gln Leu
                                105
Asp Val Ala Gly Gly Thr His Val Gly Ile Met Ala Pro Gln Ser Val
Cys Asp Ala Leu Ala Glu Ala Ile Asp His Gly Ser Glu Thr Val Leu
                        135
Asn Gly Val Pro Ala Ser Arg Leu Asn Pro Ala Gln Arg Arg Leu
                    150
                                        155
Val Leu Val Ala Pro Arg Ser Pro Glu Leu Phe Asp Asp Thr Ala Arg
                                    170
Ala Asn Ile Val Leu Asp Ser Gln Thr Thr Val Ala Arg Leu Asn Ala
                                185
                                                     190
<210> 65
<211> 339
<212> DNA
<213> Homo sapiens
<400> 65
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aagaaggccg ctttcaagat cacccgcgcc gggcaactag tgggcaccat ggcctccgag
egeettggeg taecettegg catcategae etttegettg eecetaetge egaattggga
gattegggg cecacatest tgageatatg ggattggace aagtaggeac geaeggeaca
actgctgctt tggctctgct taacgacgcc gtaaagaaag gcggcatgat ggcctgcccc
egegteggeg gtttgtetgg etectteate eegggetee
339
<210> 66
<211> 113
<212> PRT
<213> Homo sapiens
<400> 66
Val Asp Arg Ala Leu Gly Ser Leu Glu Gly Ala Ser Leu Asp Gln Val
Ala Glu Glu Val Lys Lys Ala Ala Phe Lys Ile Thr Arg Ala Gly Gln
Leu Val Gly Thr Met Ala Ser Glu Arg Leu Gly Val Pro Phe Gly Ile
Ile Asp Leu Ser Leu Ala Pro Thr Ala Glu Leu Gly Asp Ser Gly Ala
His Ile Leu Glu His Met Gly Leu Asp Gln Val Gly Thr His Gly Thr
                    70
Thr Ala Ala Leu Ala Leu Leu Asn Asp Ala Val Lys Lys Gly Gly Met
                                    90
Met Ala Cys Pro Arg Val Gly Gly Leu Ser Gly Ser Phe Ile Pro Gly
           100
                                105
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Ser
<210> 67
  <211> 446
  <212> DNA
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 Gly Ser Phe Asp Ala Asn Glu Leu Ala Val Thr Pro Asp Thr Asp Thr
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 Val Ile Gln Gly Val Gly Pro Ala Leu Ala Leu Leu Asp Ser Ala Trp
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                                              60
 Gly Arg Gln Ile His Val Glu Thr Thr Gly Cys Pro Ser Ala Val Val
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                                          75
 Trp Asn Pro Arg Ser Ser Thr His Ala Asp Asn Pro Thr Ala Gln
 Ala Trp Arg Asp Phe Val Cys Val Glu Thr Gly Ala Cys Lys Asp Asn
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 Ala Val Ile Val Ala Pro His Ser Asp Leu Thr Met Ser Thr Arg Ile
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453

170

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Pro Arg Leu His Leu Ser Cys Thr 180

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Phe Ile Leu Pro Asn Ala Leu Ile Arg Thr Ala Gly Gly Arg Glu Val
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Arg Gly Val Trp Cys Asn Asn Leu Val Tyr Asn Ile His Leu Leu Thr
Gly Lys Ile Ser Thr Pro Gly Asn Ser Pro Phe Ser Leu Thr Gly Gln
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Pro Ser Ala Cys Gly Thr Ala Arg Glu Val Gly Thr Phe Ser His Arg
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Leu Pro Ala Asp Met Val Val Thr Ser Lys Ala His Arg Asp Ile Ala
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Arg 545 Asp Thr Glu Ser Gly 625 Asp	530 Glu Tyr His Val Gly 610 Cys	Cys Tyr Ser Glu 595 Pro Asp Val	Val Glu Pro 580 Arg Ala His	Val Pro 565 Leu Ala Val Asp Tyr 645	Gly 550 Ala Ala Pro Ala Cys 630 Ala	Phe Arg Ala Pro 615 Gly Ser	Thr Glu Glu Arg 600 Glu Ala Ala	Ser Ala Leu 585 Gly Glu Gln Cys	Ala Thr 570 Cys Pro Gly Gly Arg 650	Leu 555 Arg Ala Gly Ala Asn 635 Leu	Phe Gly Trp Ala 620 Pro	Val Tyr Pro Phe 605 Ile Val Glu	Ser Asn Ala 590 Pro Ala Cys Ala	Val 575 Cys Gly Arg Gly Ala 655	Tyr 560 Ser Asn Glu Cys Ser 640 Cys
Arg 545 Asp Thr Glu Ser Gly 625 Asp	530 Glu Tyr His Val Gly 610 Cys	Cys Tyr Ser Glu 595 Pro Asp Val	Val Glu Pro 580 Arg Ala His Val	Val Pro 565 Leu Ala Val Asp Tyr 645	Gly 550 Ala Ala Pro Ala Cys 630	Phe Arg Ala Pro 615 Gly Ser	Thr Glu Glu Arg 600 Glu Ala Ala	Ser Ala Leu 585 Gly Glu Gln Cys Ala	Ala Thr 570 Cys Pro Gly Gly Arg 650	Leu 555 Arg Ala Gly Ala Asn 635 Leu	Phe Gly Trp Ala 620 Pro	Val Tyr Pro Phe 605 Ile Val Glu	Ser Asn Ala 590 Pro Ala Cys Ala Cys	Val 575 Cys Gly Arg Gly Ala 655	Tyr 560 Ser Asn Glu Cys Ser 640 Cys
Arg 545 Asp Thr Glu Ser Gly 625 Asp	530 Glu Tyr His Val Gly 610 Cys Gly Gln	Cys Tyr Ser Glu 595 Pro Asp Val Ala	Val Glu Pro 580 Arg Ala His Val Ala 660	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu	Phe Arg Ala Pro 615 Gly Ser	Thr Glu Glu Arg 600 Glu Ala Ala Pro	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro	Leu 555 Arg Ala Gly Ala Asn 635 Leu	Phe Gly Trp Ala 620 Pro Arg	Val Tyr Pro Phe 605 Ile Val Glu Cys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670	Val 575 Cys Gly Arg Gly Ala 655 Ala	Tyr 560 Ser Asn Glu Cys Ser 640 Cys
Arg 545 Asp Thr Glu Ser Gly 625 Asp	530 Glu Tyr His Val Gly 610 Cys Gly Gln	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg	Val Glu Pro 580 Arg Ala His Val Ala 660	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala	Phe Arg Ala Pro 615 Gly Ser	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro	Leu 555 Arg Ala Gly Ala Asn 635 Leu	Phe Gly Trp Ala 620 Pro Arg	Val Tyr Pro Phe 605 Ile Val Glu Cys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670	Val 575 Cys Gly Arg Gly Ala 655 Ala	Tyr 560 Ser Asn Glu Cys Ser 640 Cys
Arg 545 Asp Thr Glu Ser Gly 625 Asp Arg	530 Glu Tyr His Val Gly 610 Cys Gly Gln	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu	Phe Arg Ala Pro 615 Gly Ser Glu Ser	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro	Phe Gly Trp Ala 620 Pro Arg Ser	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp	Val Val 575 Cys Gly Arg Gly Ala 655 Ala Asp	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu
Arg 545 Asp Thr Glu Ser Gly 625 Asp Arg	530 Glu Tyr His Val Gly 610 Cys Gly Gln Gln ser	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro	S40 Pro Phe Gly Trp Ala 620 Pro Arg Ser Tyr	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp	Val Val 575 Cys Gly Arg Gly Ala 655 Ala Asp	Tyr 560 Ser Asn Glu Cys Ser 640 Cys
Arg 545 Asp Thr Glu Ser Gly 625 Asp Arg Glu Ala	530 Glu Tyr His Val Gly 610 Cys Gly Gln Gln ser 690	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675 Val	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu Ala	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu Ala	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro 695	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680 Leu	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser Gln	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser Gln	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro Thr	S40 Pro Phe Gly Trp Ala 620 Pro Arg Ser Tyr Val 700	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685 Lys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp Leu	Val 575 Cys Gly Arg Gly Ala 655 Ala Asp	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu Gly
Arg 545 Asp Thr Glu Ser Gly 625 Asp Glu Ala Ala	530 Glu Tyr His Val Gly 610 Cys Gly Gln Gln ser 690	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675 Val	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu Ala	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro Pro	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu Ala Gly Glu	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro 695	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680 Leu	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser Gln	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser Gln	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro Thr Asp	S40 Pro Phe Gly Trp Ala 620 Pro Arg Ser Tyr Val 700	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685 Lys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp Leu	Val 575 Cys Gly Arg Gly Ala 655 Ala Asp	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu Gly Ala
Arg 545 Asp Thr Glu Ser Gly 625 Asp Glu Ala Ala 705	530 Glu Tyr His Val Gly 610 Cys Gly Gln Gln Ser 690 Gly	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675 Val Leu	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu Ala Glu	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro Pro Val	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu Ala Gly Glu 710	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro 695 Asp	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680 Leu Ser	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser Gln Asp	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser Gln Pro	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro Thr Asp Glu 715	Phe Gly Trp Ala 620 Pro Arg Ser Tyr Val 700 Pro	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685 Lys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp Leu Gly	Val Val 575 Cys Gly Arg Gly Ala 655 Ala Asp Asn Glu	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu Gly Ala 720
Arg 545 Asp Thr Glu Ser Gly 625 Asp Glu Ala Ala 705	530 Glu Tyr His Val Gly 610 Cys Gly Gln Gln Ser 690 Gly	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675 Val Leu	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu Ala Glu	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro Pro Val Thr	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu Ala Gly Glu	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro 695 Asp	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680 Leu Ser	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser Gln Asp	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser Gln Pro	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro Thr Asp Glu 715	Phe Gly Trp Ala 620 Pro Arg Ser Tyr Val 700 Pro	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685 Lys	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp Leu Gly	Val Val 575 Cys Gly Arg Gly Ala 655 Ala Asp Asn Glu Gly	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu Gly Ala 720
Arg 545 Asp Thr Glu Ser Gly 625 Asp Arg Glu Ala 705 Glu	530 Glu Tyr His Val Gly 610 Cys Gly Gln Ser 690 Gly Asp	Cys Tyr Ser Glu 595 Pro Asp Val Ala Arg 675 Val Leu Arg	Val Glu Pro 580 Arg Ala His Val Ala 660 Leu Ala Glu Val	Val Pro 565 Leu Ala Val Asp Tyr 645 Pro Pro Val Thr 725	Gly 550 Ala Ala Pro Ala Cys 630 Ala Leu Ala Gly Glu 710	Phe Arg Ala Pro 615 Gly Ser Glu Ser Pro 695 Asp Gly	Thr Glu Glu Arg 600 Glu Ala Ala Pro Ser 680 Leu Ser	Ser Ala Leu 585 Gly Glu Gln Cys Ala 665 Ser Gln Asp Arg	Ala Thr 570 Cys Pro Gly Gly Arg 650 Pro Ser Gln Pro 730	Leu 555 Arg Ala Gly Ala Asn 635 Leu Pro Thr Asp Glu 715 Pro	Phe Gly Trp Ala 620 Pro Arg Ser Tyr Val 700 Pro	Val Tyr Pro Phe 605 Ile Val Glu Cys Gly 685 Lys Glu Ser	Ser Asn Ala 590 Pro Ala Cys Ala Cys 670 Asp Leu Gly Ser	Val Val 575 Cys Gly Arg Gly Ala 655 Ala Asp Asn Glu Gly 735	Tyr 560 Ser Asn Glu Cys Ser 640 Cys Leu Leu Gly Ala 720 Asn

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745
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Thr Pro Ala Pro Gln Arg His Ser Gly Arg Val Val Gly Ala His Arg
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Pro Gly Leu Leu Ser Pro Val Phe Val Tyr Ser Pro Ala Phe Gln Ser
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Lys Ala Glu Tyr Leu Ala Ala Thr Ser Tyr Arg Asp Phe Leu Leu Lys
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Asp Val Gly Leu Ser Glu Gly Ala Val Lys Tyr Phe Gln Ser Arg Thr
Asn Asp Phe Met Ala Leu Ser Ile Asp Ala Val Ala Ser Ala Asp Ala
Tyr Ser Val Gly Phe Pro Gly Phe Gly Gly Met Asn Leu Ala Pro Ile
Ser Glu Glu Ala Ala Ala Glu Met Glu Glu Pro Tyr Ile Tyr His Phe
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Pro Asp Gly Asn Ala
            100
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cctctatgag tggaatccag tctcatggcc ccccccatgg ctcctgttac cctggaggag
getactecat gaggetgget eeggtggeae tgeetegggg etgeatgtae atgtgtgtge
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cgtgcatgtg tgtgcacgtg tgtgcatgtg cacgtgtgcg cacccgtgtg catgtccgtg
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Met Xaa Val Cys Met Cys Val Cys Thr Cys Xaa Cys Val Pro Val Cys
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Met Cys Pro Cys Ala Cys Leu Cys Cys Val Cys Ala Cys Met Cys Ala
Cys Leu Cys Val Xaa Val Cys Val Arg Ala Cys Val Cys Thr Cys Val
His Val His Val Cys Ala Pro Val Cys Met Ser Val Cys Thr Arg
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Gly Leu Thr Lys Val Gln Lys Ile Val Ala Ala Ser Glu Phe Leu Arg
                                25
Asn Asp Leu Ile Gly Leu Gly Ile Asp Lys Ala Lys Ile Glu Ile Ile
His Asn Gly Ile Asp His Arg Pro Phe Phe Pro Gln Leu Gln Ile Asp
Ala Glu Thr Val Thr Ile Lys Pro Phe Ala Ile Lys Arg Pro Tyr Phe
                    70
Ile Tyr Gly Ser Arg Leu Ser Gly Pro Glu Lys Lys His Ile Glu Leu
Ile Lys Ala Phe Ala Leu Phe Lys Glu Arg Thr Lys Ser Pro His Pro
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Leu Val Ile Ala Gly Ala Glu Gly Pro Ser Ser Glu Glu Val His
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Asp Lys Ala Arg Ile Leu Asp Ala Val Lys Leu Leu Ser Ser Leu Gly
Phe Lys Val Ile Ala Thr Ser Gly Thr Gln Arg Phe Leu Val Glu Asn
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## ATTORNEY DOCKET NO.: 15966-543

<210> 87

<211> 355

<212> DNA

<213> Homo sapiens

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attgtgcttc gcccggtgac gagtgaggac gccatgactg cggactgggc acgtatccca 180

tatgacgtac tggaaaagat ctcgactcgc attacgaatg cgtgtccgca aatcaaccgg 240

gtggtactcg atatcacatc taaaccgccg gccaccatcg agtgggaatg agccccgtct 300

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<210> 88

<211> 96

<212> PRT

<213> Homo sapiens

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Cys Pro Val Val Leu Leu Ser Asp Val His Ser Val Gly Val Gln Gly

Cys Pro Val Val Leu Leu Ser Asp Val His Ser Val Giy Val Gin Giy
20 25 30

Asp Gly Arg Thr Tyr Gly Ser Pro Ile Val Leu Arg Pro Val Thr Ser 35 40 45

Glu Asp Ala Met Thr Ala Asp Trp Ala Arg Ile Pro Tyr Asp Val Leu 50 55 60

Glu Lys Ile Ser Thr Arg Ile Thr Asn Ala Cys Pro Gln Ile Asn Arg
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Val Val Leu Asp Ile Thr Ser Lys Pro Pro Ala Thr Ile Glu Trp Glu 85 90 95

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ATTORNEY DOCKET NO.: 15966-543

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## ATTORNEY DOCKET NO.: 15966-543

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90

Ala His Val Arg Arg Pro Ile Arg Ala Cys Cys Ser Gly Met Gly Pro

His Arg Asn Ala Pro Arg Gly Thr Ala Cys Arg Thr Arg

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Pro Phe Pro Ser Ser Asp Ile Thr Pro Asn Ile Glu Arg Leu Ser Asn
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Ser Pro Leu Arg Ser Ser Arg Ala Tyr Ala Gln Asp Phe Met Thr Leu
                            40
Asp Lys Arg Trp His Val Met Ala Gly Ile Ala Leu Asn Gln Leu Pro
Gln Glu Gly Gly Pro Thr Glu Arg Ala Trp Thr Pro Lys Leu Gly Leu
Ser Phe Asp Val Ser Asp Thr Met Ser Leu Tyr Gly Ala Tyr Ser Arg
Gly Phe Ser Thr Tyr Gln Pro Ala Arg Lys Ala Pro Arg Ala Tyr Gly
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Pro Ser Ala Ala Arg Pro Ser Lys Arg Glu
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240
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Arg Pro Arg Arg Trp Ala Pro Lys Ile Thr Ala Ala Ser Pro Ala Ser
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Arg Thr Arg Asp Ser Ser Lys Ser Arg Val Met Gly Ser Thr Ile Arg
Ser Ala Trp Ser Met Arg Asn Ser Arg Gly Arg Leu Leu Gly Arg Arg
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Gly Arg Trp Val Ser Thr Val Ile Ala Glu Arg Ser Ser Ser Thr Thr
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 Ser Gly Ala Asp Ala
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619
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Cys Ala Ser Arg Asn Val Thr Ala Cys Leu His Pro His Trp Gly Thr
Met Val Gln Asn Arg Asp Glu Val Ile Arg Val Leu Glu Asn Ser Ser
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Ile Gly Leu Cys Leu Asp Thr Gly His Leu Ala Cys Gly Gly Thr Asp
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Val Val Glu Leu Val Arg Lys Tyr Ala Asn Arg Val Asp Ile Val His
                                105
Ala Lys Asp Val His Lys Glu Met Ala Asp Lys Leu Leu Pro Gly Glu
        115
Ile Thr Trp Ser Glu Gly Ile Arg Ala Gly Met Phe Ala Pro Ile Gly
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Gly Gly Asp Pro Ser His Trp Gly Glu Thr Pro Ala Met Gly Lys Asp
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Pro Cys His Trp Gly Arg Xaa Pro Ala Ile Gly Gly Asp Pro Cys Arg
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Ser Pro His Cys Phe Arg Leu Leu Arg Pro Val Leu Val Thr Asp Arg
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ctcattacgg ctgcgacggc gggcgcctgt ctcggttttt tgccccacaa ctggcatccg
gcgaggatgt tcatgggtga ttccggagct ctgctacttg gcttattgct a
471
<210> 118
<211> 157
<212> PRT
<213> Homo sapiens
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<400> 118
Xaa Ala Leu Thr Ile Cys Leu Ala Gly Val Val Ile Cys Ala Val Gly
Val Val Asp Asp Leu Leu Asp Leu Pro Ala Leu Ala Lys Ala Ala Gly
Gln Val Leu Ala Ala Gly Ile Val Val Thr Gly Gly Val Arg Met Phe
                            40
Trp Ile Pro Leu Pro Asn Ser Ile Ile Ala Leu Gly Thr Pro Thr Ser
                                            60
                        55
Ile Leu Val Thr Val Phe Phe Ile Val Leu Cys Ala Asn Ala Val Asn
Phe Ile Asp Gly Leu Asp Gly Leu Ala Ser Gly Val Val Ala Ile Gly
Ser Leu Ala Phe Phe Ser Tyr Thr Tyr Leu Leu Ala His Glu Gln Asp
                                105
            100
Phe Val Val Ala Thr Thr Ser Leu Ile Thr Ala Ala Thr Ala Gly
                                                125
                            120
        115
Ala Cys Leu Gly Phe Leu Pro His Asn Trp His Pro Ala Arg Met Phe
                        135
Met Gly Asp Ser Gly Ala Leu Leu Leu Gly Leu Leu Leu
145
<210> 119
<211> 302
<212> DNA
<213> Homo sapiens
<400> 119
ntcaaacatg agcagtcgtg gcggccgagg ccgcggtggc tattatcgcg agctttatgg
tagecgaggt cgaggcagta aatetaatga aaetttegea aaaaattegg atgtetaete
tcagaaaaag actcgaacag tacgaggcac ctccgaagat ttagcacgat cgctccataa
getteatatg egecegtace etgegtatea tgaeattgag ggtatgtggg ettteecage
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cn
302
<210> 120
 <211> 98
 <212> PRT
 <213> Homo sapiens
 <400> 120
Met Ser Ser Arg Gly Gly Arg Gly Arg Gly Gly Tyr Tyr Arg Glu Leu
Tyr Gly Ser Arg Gly Arg Gly Ser Lys Ser Asn Glu Thr Phe Ala Lys
                                 25
 Asn Ser Asp Val Tyr Ser Gln Lys Lys Thr Arg Thr Val Arg Gly Thr
 Ser Glu Asp Leu Ala Arg Ser Leu His Lys Leu His Met Arg Pro Tyr
```

```
60
                        55
   50
Pro Ala Tyr His Asp Ile Glu Gly Met Trp Ala Phe Pro Ala Phe Thr
                                        75
                    70
Phe Tyr Leu Asp His Ala Gln Ala Asp Pro Tyr Ala Ala Pro Asn Lys
                                    90
Ala Arg
<210> 121
<211> 318
<212> DNA
<213> Homo sapiens
<400> 121
ngcatggggg gccctgggac cgcacttgtg cccctttttt ttttagggaa aaaattgagc
cctaaaggat ttgccgcatt acaggaaagt tttttggtaa gtttggggtt gtttctgtgc
tgtgtgagaa ggagtagaag cagctccagt agagtgggcc ttttcatttt tatccagagg
aaatttgtag getgtggeta ttaetteett ttttttett ttttttttg tttagagaca
gagtetgnet etgtegecag getggagtga agtggeacga teteagetea etgeaacete
300
tgcctcccag gttcaagc
318
<210> 122
<211> 89
<212> PRT
<213> Homo sapiens
<400> 122
Xaa Met Gly Gly Pro Gly Thr Ala Leu Val Pro Leu Phe Phe Leu Gly
                                     10
Lys Lys Leu Ser Pro Lys Gly Phe Ala Ala Leu Gln Glu Ser Phe Leu
Val Ser Leu Gly Leu Phe Leu Cys Cys Val Arg Arg Ser Arg Ser Ser
                             40
Ser Ser Arg Val Gly Leu Phe Ile Phe Ile Gln Arg Lys Phe Val Gly
                                             60
                        55
Cys Gly Tyr Tyr Phe Leu Phe Phe Leu Phe Phe Phe Cys Leu Glu Thr
                                         75
                    70
Glu Ser Xaa Ser Val Ala Arg Leu Glu
                85
<210> 123
<211> 338
<212> DNA
<213> Homo sapiens
<400> 123
acgcgtctag ggtagaaatc aactccagta actgtcattc aacctcagca atgctgggga
60
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cgggcagagg cagggcagct gtgtgccaca ttcctgccag ggctggtcag gccccggctc
tcaccactcc tcctccctgc tttgaacctg tggaacaaag ggcccctgca ccccaactca
ttcctctttg ccacataagg gcctcaagtc atgctgtccc ctctgcctgg gttgcttttt
ctccctctgc ttgggtcact gttcacacca ctggccactt tcctcaggga agggccctca
ctgcccacac acctaaacat gccccctgct cctccata
338
<210> 124
<211> 96
<212> PRT
<213> Homo sapiens
<400> 124
Met Leu Gly Thr Gly Arg Gly Arg Ala Ala Val Cys His Ile Pro Ala
Arg Ala Gly Gln Ala Pro Ala Leu Thr Thr Pro Pro Pro Cys Phe Glu
                                                     30
Pro Val Glu Gln Arg Ala Pro Ala Pro Gln Leu Ile Pro Leu Cys His
Ile Arg Ala Ser Ser His Ala Val Pro Ser Ala Trp Val Ala Phe Ser
Pro Ser Ala Trp Val Thr Val His Thr Thr Gly His Phe Pro Gln Gly
                                        75
                    70
Arg Ala Leu Thr Ala His Thr Pro Lys His Ala Pro Cys Ser Ser Ile
                85
                                    90
<210> 125
<211> 280
<212> DNA
<213> Homo sapiens
<400> 125
ccatggacct ggccagccac catcacctgc ctcctgcctc acccaccctg ggtgcctgcc
ggcaaggatt ggagggcaga ctgctggagc gtgagaccag gccaatctgt ctttctggga
accttcagcc tccaactgga gctgactgtc aactttcggg tgagaagtca cttttctgca
ttcccaccac actatctatc tgtgcaatac ggcagcgtga cagcactcac cttattgagg
gettetgetg teetggeeca ttetggatag geetgateta
280
<210> 126
<211> 92
<212> PRT
<213> Homo sapiens
<400> 126
Met Asp Leu Ala Ser His His His Leu Pro Pro Ala Ser Pro Thr Leu
```

```
10
 1
Gly Ala Cys Arg Gln Gly Leu Glu Gly Arg Leu Leu Glu Arg Glu Thr
                              25
Arg Pro Ile Cys Leu Ser Gly Asn Leu Gln Pro Pro Thr Gly Ala Asp
Cys Gln Leu Ser Gly Glu Lys Ser Leu Phe Cys Ile Pro Thr Thr Leu
Ser Ile Cys Ala Ile Arg Gln Arg Asp Ser Thr His Leu Ile Glu Gly
Phe Cys Cys Pro Gly Pro Phe Trp Ile Gly Leu Ile
               85
<210> 127
<211> 444
<212> DNA
<213> Homo sapiens
<400> 127
egegtgateg eegtggegga gggeegegge geegaetega tegeeeaget gaeaacegag
ctgcaaagcc gtcactgccc tgcggagcag atcacgtccg tcagcatcga catgtcgcca
gegttcatca ggggctgcgc cgagcacctg cccaacgcgc gcgtcacctt cgacaagttc
gacaagteee teaaggggat gegetggteg etgetgaaga acegegeeag eeteaageee
gaggetgeeg cegatetgga tgeectgate geeaggatgg ceaetgtgeg caeeggege
gcctgggtct acaaggagca gctgcgcgag atcctcgcgc gcaagcagat caacgtggca
cgcgacatgc tcaagcactg gtgc
444
<210> 128
<211> 148
<212> PRT
<213> Homo sapiens
<400> 128
Arg Val Ile Ala Val Ala Glu Gly Arg Gly Ala Asp Ser Ile Ala Gln
Leu Thr Thr Glu Leu Gln Ser Arg His Cys Pro Ala Glu Gln Ile Thr
Ser Val Ser Ile Asp Met Ser Pro Ala Phe Ile Arg Gly Cys Ala Glu
                            40
His Leu Pro Asn Ala Arg Val Thr Phe Asp Lys Phe His Val Ile Gly
                        55
His Ala Asn Ala Ala Val Asp Arg Met Arg Arg Ile Glu Gln Arg Ser
                                       75
Asp Lys Ser Leu Lys Gly Met Arg Trp Ser Leu Leu Lys Asn Arg Ala
 Ser Leu Lys Pro Glu Ala Ala Ala Asp Leu Asp Ala Leu Ile Ala Arg
```

```
105
            100
Met Ala Thr Val Arg Thr Ala Arg Ala Trp Val Tyr Lys Glu Gln Leu
                            120
Arg Glu Ile Leu Ala Arg Lys Gln Ile Asn Val Ala Arg Asp Met Leu
   130
                        135
Lys His Trp Cys
145
<210> 129
<211> 291
<212> DNA
<213> Homo sapiens
<400> 129
gaggagggac gtaccgtccc cgttatagcc aagctcgaga agccgcaagc tatcgagaac
ttggacgaga ttattgacgt ctttgacgcc gtcatggttg cccgtggcga tatggccgtc
120
gagtgcccgc tcgaggaagt tccgctgatc caaaagcaga tcatcgagaa ggctcgttta
caggetaage cegteattgt ggecacecag atgettgagt egatgateca egeteeeegt
cegaceegeg etgaggeege egacgtegeg aaegeeatee ttgaeggege g
291
<210> 130
<211> 97
<212> PRT
<213> Homo sapiens
<400> 130
Glu Glu Gly Arg Thr Val Pro Val Ile Ala Lys Leu Glu Lys Pro Gln
Ala Ile Glu Asn Leu Asp Glu Ile Ile Asp Val Phe Asp Ala Val Met
                                25
Val Ala Arg Gly Asp Met Ala Val Glu Cys Pro Leu Glu Glu Val Pro
Leu Ile Gln Lys Gln Ile Ile Glu Lys Ala Arg Leu Gln Ala Lys Pro
                        55
Val Ile Val Ala Thr Gln Met Leu Glu Ser Met Ile His Ala Pro Arg
Pro Thr Arg Ala Glu Ala Ala Asp Val Ala Asn Ala Ile Leu Asp Gly
Ala
<210> 131
<211> 416
<212> DNA
<213> Homo sapiens
<400> 131
teeggagegt eegtggeeet catgggtgtg teagegtggt tgetgteteg ggeegeagag
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attecacegg tgetetacet ggaggeegea geegtegggg ttegattett eggeatetee
cgcggtgtct tccgctacgc cgaacgtctg gtaggccacg acctggctct gcggatgcag
ggggcattgc ggatgcgggt ctacgaccgg ctgtcacgta ccnaccctgc tgggnnacgt
cgccggggtg acctgctggt acgggttact gccgacgtcg acgcggtgtt ggacatggtc
gtgegggtga tegtteegge gtgegegtea ageetegtea teattggeae caeggteett
ctttgtccga gagaaggttg agttttctta gccggattcc aacacagcct gggggc
<210> 132
<211> 126
<212> PRT
<213> Homo sapiens
<400> 132
Ser Gly Ala Ser Val Ala Leu Met Gly Val Ser Ala Trp Leu Leu Ser
Arg Ala Ala Glu Ile Pro Pro Val Leu Tyr Leu Glu Ala Ala Ala Val
Gly Val Arg Phe Phe Gly Ile Ser Arg Gly Val Phe Arg Tyr Ala Glu
Arg Leu Val Gly His Asp Leu Ala Leu Arg Met Gln Gly Ala Leu Arg
                        55
Met Arg Val Tyr Asp Arg Leu Ser Arg Thr Xaa Pro Ala Gly Xaa Arg
                                         75
                    70
Arg Arg Gly Asp Leu Leu Val Arg Val Thr Ala Asp Val Asp Ala Val
                                     90
Leu Asp Met Val Val Arg Val Ile Val Pro Ala Cys Ala Ser Ser Leu
                                 105
            100
Val Ile Ile Gly Thr Thr Val Leu Leu Cys Pro Arg Glu Gly
                             120
        115
<210> 133
<211> 327
<212> DNA
<213> Homo sapiens
<400> 133
geegttgeta tegetgetgg tatgegtgea gaegteactg tittigatat caatateget
gegttgaaga gactegeega catetaceag ggtegtgtte acaeagtagt atecaeeege
geegaaattg egaaggeget agaaaceget gaegttgtga teggttetgt eettatteeg
ggtagttcta ccccgaagct tgttactacc gatatggttg ctcacatgca gcctgggtct
gttcttattg atattgctat agaccaaggc ggctgcttcg aggattcgca ccccaccact
 tacgatgacc ccactttcac tgtgcac
 327
```

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<210> 134
<211> 109
<212> PRT
<213> Homo sapiens
<400> 134
Ala Val Ala Ile Ala Ala Gly Met Arg Ala Asp Val Thr Val Phe Asp
Ile Asn Ile Ala Ala Leu Lys Arg Leu Ala Asp Ile Tyr Gln Gly Arg
                                25
Val His Thr Val Val Ser Thr Arg Ala Glu Ile Ala Lys Ala Leu Glu
Thr Ala Asp Val Val Ile Gly Ser Val Leu Ile Pro Gly Ser Ser Thr
                        55
Pro Lys Leu Val Thr Asp Met Val Ala His Met Gln Pro Gly Ser
                                        75
Val Leu Ile Asp Ile Ala Ile Asp Gln Gly Gly Cys Phe Glu Asp Ser
                                                         95
                                    90
His Pro Thr Thr Tyr Asp Asp Pro Thr Phe Thr Val His
<210> 135
<211> 560
<212> DNA
<213> Homo sapiens
<400> 135
taagatgtgg teetgeeetg tteetgaagg ggetgeaget etgatggaaa atacagggat
ttacactcag ggctacagcc acggggggct gaggcccaag gctgcaatct cgggggaagg
120
ggaagttggc ttttcctggt ggattggaaa catcctcttg gaggcaaaga cttttcctgg
atcttacaga cttcccggga tttttagatt agaatattgg gggcaaagga ggctgtcttg
ttttaaagca atgctacata gacacagtgg ggaagacctg gttcgacggc agataagcag
tgggtgatgg gcttgaggag gagagtcagg gcaaagtcta agactgagca gaaaggaatt
cocccatctc ccatggataa gtacgttcta gaacattctc tttgggtcta atactctgaa
420
atgacatett gtetteatge tegagagaga attactteae tggeteeaet tggagtgeea
gtgttcagac accaagcctg actgggaggg ttccgttttc ttaacacctt cccaccgccg
 540
 acttccaagt ccccacgcgt
 560
 <210> 136
 <211> 100
 <212> PRT
 <213> Homo sapiens
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<400> 136
Met Trp Ser Cys Pro Val Pro Glu Gly Ala Ala Ala Leu Met Glu Asn
Thr Gly Ile Tyr Thr Gln Gly Tyr Ser His Gly Gly Leu Arg Pro Lys
Ala Ala Ile Ser Gly Glu Gly Glu Val Gly Phe Ser Trp Trp Ile Gly
                            40
Asn Ile Leu Leu Glu Ala Lys Thr Phe Pro Gly Ser Tyr Arg Leu Pro
                        55
Gly Ile Phe Arg Leu Glu Tyr Trp Gly Gln Arg Arg Leu Ser Cys Phe
Lys Ala Met Leu His Arg His Ser Gly Glu Asp Leu Val Arg Arg Gln
                                    90
Ile Ser Ser Gly
            100
<210> 137
<211> 429
<212> DNA
<213> Homo sapiens
<400> 137
accggttgga tggcctgcag gccaaagcgt tcctgcaaac tcagcaggcc ttcagcgcaa
gaggcaaaca gctggtcgcg cacctgcttg aggtccaccg attgcgcatc gcccttgagc
aaggegegee agttggtttt gteggeeact tggetgegga acaggtette gacaaaaceg
gactgctggc gggtcgcaac gcgcatgatc ggcagcgcct ggctggcgcc ctggtcgagc
cagegegteg geagttgggt ggeeegggtg atacegaeet tgateeeega egaattggee
aggtacacca catggtcggt catgcagaat gtttcgcccc agccgggatc acggcaagtg
ccggcgtcgt aatggcaacg ttcggggctc atgatgcaca ggtcacactg ggccagcttg
420
gtcatgccc
429
<210> 138
<211> 141
<212> PRT
<213> Homo sapiens
<400> 138
Met Thr Lys Leu Ala Gln Cys Asp Leu Cys Ile Met Ser Pro Glu Arg
Cys His Tyr Asp Ala Gly Thr Cys Arg Asp Pro Gly Trp Gly Glu Thr
Phe Cys Met Thr Asp His Val Val Tyr Leu Ala Asn Ser Ser Gly Ile
                            40
Lys Val Gly Ile Thr Arg Ala Thr Gln Leu Pro Thr Arg Trp Leu Asp
                        55
Gln Gly Ala Ser Gln Ala Leu Pro Ile Met Arg Val Ala Thr Arg Gln
```

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65
                    70
Gln Ser Gly Phe Val Glu Asp Leu Phe Arg Ser Gln Val Ala Asp Lys
                                    90
Thr Asn Trp Arg Ala Leu Leu Lys Gly Asp Ala Gln Ser Val Asp Leu
                                105
Lys Gln Val Arg Asp Gln Leu Phe Ala Ser Cys Ala Glu Gly Leu Leu
                           120
Ser Leu Gln Glu Arg Phe Gly Leu Gln Ala Ile Gln Pro
                        135
   130
<210> 139
<211> 341
<212> DNA
<213> Homo sapiens
<400> 139
acgcgtcgtt tgaaggcttg atccgcacgt ccaattcgct ttgcgccaat gcgccgcagc
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tttgcagcct gtaacgactg agggttcgga tggaaaaaca catgctccag gatgggaccg
acggccactt caccgatctc ttcatagccc tggcgtttgt agaaatccag gtagcgcgaa
tegecagegt egageaegae geetgatgag tgegggteat t
<210> 140
<211> 113
<212> PRT
<213> Homo sapiens
Met Thr Arg Thr His Gln Ala Ser Cys Ser Thr Leu Ala Ile Arg Ala
Thr Trp Ile Ser Thr Asn Ala Arg Ala Met Lys Arg Ser Val Lys Trp
                                25
Pro Ser Val Pro Ser Trp Ser Met Cys Phe Ser Ile Arg Thr Leu Ser
Arg Tyr Arg Leu Gln Arg Phe Glu Thr Glu Leu Phe Arg Gln Phe Arg
Val Gln Ser Val Ser Pro Ala Arg Val Ala Ser Pro Pro Met Lys Leu
                    70
Pro Gly Arg Phe Thr Ser Gly Leu Ile Leu Leu Phe Thr Ser Cys Gly
                                    90
Ala Leu Ala Gln Ser Glu Leu Asp Val Arg Ile Lys Pro Ser Asn Asp
                                105
Ala
<210> 141
<211> 324
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<212> DNA
<213> Homo sapiens
<400> 141
gaatteetet tggatagett egggtaaatg ggtacageaa atateaggag egcaacegea
acctttactt actggtacat gaacaccatt tacattacag ctatcgtact caccccacgt
catgtgaaca gacacataac tgaaaggttt ataaaccaca gtctcacggt acgtatgacc
gtcaactgtg aacaccgcta agtaatagcc tgcgggggct tgcatgaact cctttgacca
tgcgtaataa atacgtccgt cattagtcac acctgatggg gcgaaacaaa aagaacggca
gcagttatca ccgcccatac gcgt
324
<210> 142
<211> 106
<212> PRT
<213> Homo sapiens
<400> 142
Met Gly Gly Asp Asn Cys Cys Arg Ser Phe Cys Phe Ala Pro Ser Gly
                                    10
Val Thr Asn Asp Gly Arg Ile Tyr Tyr Ala Trp Ser Lys Glu Phe Met
Gln Ala Pro Ala Gly Tyr Tyr Leu Ala Val Phe Thr Val Asp Gly His
Thr Tyr Arg Glu Thr Val Val Tyr Lys Pro Phe Ser Tyr Val Ser Val
His Met Thr Trp Gly Glu Tyr Asp Ser Cys Asn Val Asn Gly Val His
                                         75
Val Pro Val Ser Lys Gly Cys Gly Cys Ala Pro Asp Ile Cys Cys Thr
His Leu Pro Glu Ala Ile Gln Glu Glu Phe
                                 105
            100
<210> 143
<211> 1325
<212> DNA
<213> Homo sapiens
<400> 143
nacgcgtgga tctgccagct gagcctggag ctgtgcaggc agctgccctg ctacgatgag
gcaccccagg agaagaactt cctgtacaaa tgcataggca ccaccctggg tgctgcttca
agtaaggagg tggtgaggaa gcaccttcaa gagctgctgg agacggccag ataccaggag
gaggcagaac gcgagggcct cgcctgctgc ttcgggatct gtgccatctc ccacctcgag
gacacgctgg cccagctgga ggacttcgtg aggtcagagg tcttcagaaa atccattggc
300
```

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attctcaaca tttttaagga tcgaagtgag aacgaagtgg agaaggtgaa gagtgctctg
atcctgtgct atgggcacgt ggcggcccgg gccccccggg agctggtgct ggccaaggta
gagtcagaca tcctccggaa catcntgcca gcacttcagc acnncaagga cccagccctg
aagetgtgee ttgteeagag tgtgtgeatg gteageegeg ceatetgeag cageacceag
gctggctcct tccacttcac ccggaaagca gagctggtgg cacagatgat ggagttcatc
agggcagage ecceggacte ettgaggaca ectattegga agaaageeat geteacetge
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ggotgcotgc acagcatcat ggocotgctg cotgagocca aggaggagga cggaggotgc
cagaagteee tgtatetgga gacactgeae gecettgagg atetgetgae gageeteetg
cageggaaca tgaccecca aggeetgeag ateatgattg ageacetgag eccatggate
aagteeccaa gaggteaegt ageggegegt geectaggee tgagegeeet eetegtgege
tacttectgg agcacetgeg tgteagtgge geceaagtag ataceaggtt tecatetgag
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1080
cttctcccca cggtgtgcgg acctgtggcc tgccacccgc caggaggccg tggactgtgt
ctactccctg ctgtacctcc agctcggcta tgagggcttc tcccgggact accgcgatga
cgtggcggag cggctcctca gcctcaagga cggcctcgtg caccctgacc ccgccattct
cttccacacc tgccacagtg taggccagat tattgccaag cgcctcccc cagcccttca
1320
cgcgt
1325
<210> 144
<211> 390
<212> PRT
<213> Homo sapiens
<400> 144
Xaa Ala Trp Ile Cys Gln Leu Ser Leu Glu Leu Cys Arg Gln Leu Pro
Cys Tyr Asp Glu Ala Pro Gln Glu Lys Asn Phe Leu Tyr Lys Cys Ile
                                25
Gly Thr Thr Leu Gly Ala Ala Ser Ser Lys Glu Val Val Arg Lys His
        35
Leu Gln Glu Leu Leu Glu Thr Ala Arg Tyr Gln Glu Glu Ala Glu Arg
Glu Gly Leu Ala Cys Cys Phe Gly Ile Cys Ala Ile Ser His Leu Glu
Asp Thr Leu Ala Gln Leu Glu Asp Phe Val Arg Ser Glu Val Phe Arg
```

90

85

```
Lys Ser Ile Gly Ile Leu Asn Ile Phe Lys Asp Arg Ser Glu Asn Glu
           100
Val Glu Lys Val Lys Ser Ala Leu Ile Leu Cys Tyr Gly His Val Ala
                                                125
                            120
Ala Arg Ala Pro Arg Glu Leu Val Leu Ala Lys Val Glu Ser Asp Ile
                       135
Leu Arg Asn Ile Xaa Pro Ala Leu Gln His Xaa Lys Asp Pro Ala Leu
                                       155
                   150
Lys Leu Cys Leu Val Gln Ser Val Cys Met Val Ser Arg Ala Ile Cys
                                   170
Ser Ser Thr Gln Ala Gly Ser Phe His Phe Thr Arg Lys Ala Glu Leu
                                185
Val Ala Gln Met Met Glu Phe Ile Arg Ala Glu Pro Pro Asp Ser Leu
                            200
Arg Thr Pro Ile Arg Lys Lys Ala Met Leu Thr Cys Thr Tyr Leu Val
                        215
Ser Val Glu Pro Ala Leu Asp Glu Gln Ala Arg Ala Asp Val Ile His
                                       235
                    230
Gly Cys Leu His Ser Ile Met Ala Leu Leu Pro Glu Pro Lys Glu Glu
                                   250
Asp Gly Gly Cys Gln Lys Ser Leu Tyr Leu Glu Thr Leu His Ala Leu
                                265
Glu Asp Leu Leu Thr Ser Leu Leu Gln Arg Asn Met Thr Pro Gln Gly
                                                285
                            280
Leu Gln Ile Met Ile Glu His Leu Ser Pro Trp Ile Lys Ser Pro Arg
                                            300
                        295
Gly His Val Ala Ala Arg Ala Leu Gly Leu Ser Ala Leu Leu Val Arg
                    310
Tyr Phe Leu Glu His Leu Arg Val Ser Gly Ala Gln Val Asp Thr Arg
                                    330
Phe Pro Ser Glu Pro Arg Ile Leu Cys Asn Gly Pro Gly Ala Leu Pro
                                345
            340
Gln Pro Gly Pro Ser His Arg Pro Leu Leu Pro Thr Val Cys Gly Pro
                                                365
                            360
Val Ala Cys His Pro Pro Gly Gly Arg Gly Leu Cys Leu Leu Pro Ala
                        375
Val Pro Pro Ala Arg Leu
385
<210> 145
<211> 802
<212> DNA
<213> Homo sapiens
<400> 145
cggccgtcta ggtccggctc agtgcgctgt tgctcgccgt agaacacgag gctgcgcaag
cataagcaga cgtagagagt ggtcacatcc atgtcgatgg tgtgcgcgta atgaaggtct
acatcaccct ggtgaaggcc tgcaccacta gcgtcggcac catttccccg cgtcggacaa
gacatcatgo occatatott gacagaatgt otgacatgag tatgocacgo ogagoagoac
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```
cagaggacga caccgatctg gcggacgccg cccgttcatg gcgcagatac ctcatcctcg
300
teatttgtgg cgttatcgtc gctgtcctcg gactaggcat tttcgggtat cttgcgtggt
ggtcattgtg cgatcaagct gccggggtct gtcagcgtgg tgaacccgtt atgtactggt
420
gtteggtggt etetetggee atteteggae teattategg ggtettgaeg eagatetgge
tggagaagcg ctggtggcac atgcttgcca tcgtcatccc ggctgttttc atcgtcgccg
gtatcttttt ctggctcgcc gtctaagaag gggcgtcaca gattccacaa acgacacagg
tattgatete egitttateg geteetagea geegiggiea aegiategei aleaagegat
acaggactcg tcgttcgcat cgttgttgtg ctgctgggaa acaatcccag cgatctactc
ggctaccgcc agacagttca ctcacaaccc ctcacgccgg cgcagacatc aaatcccatt
ctcgatagac ggcccacacc ac
802
<210> 146
<211> 151
<212> PRT
<213> Homo sapiens
<400> 146
Met Lys Val Tyr Ile Thr Leu Val Lys Ala Cys Thr Thr Ser Val Gly
Thr Ile Ser Pro Arg Arg Thr Arg His His Ala Pro Tyr Leu Asp Arg
                                25
Met Ser Asp Met Ser Met Pro Arg Arg Ala Ala Pro Glu Asp Asp Thr
Asp Leu Ala Asp Ala Ala Arg Ser Trp Arg Arg Tyr Leu Ile Leu Val
                         55
Ile Cys Gly Val Ile Val Ala Val Leu Gly Leu Gly Ile Phe Gly Tyr
                     70
Leu Ala Trp Trp Ser Leu Cys Asp Gln Ala Ala Gly Val Cys Gln Arg
                                     90
Gly Glu Pro Val Met Tyr Trp Cys Ser Val Val Ser Leu Ala Ile Leu
                                 105
Gly Leu Ile Ile Gly Val Leu Thr Gln Ile Trp Leu Glu Lys Arg Trp
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Arg Thr Ile Pro Thr Thr Glu Trp Asp Ile Leu Ala Arg Leu Arg Pro
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Ser Ala Asp Glu Gly Lys Ser Trp Ala Pro Ile Lys Gly Pro Glu Gln
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Gly Gln Ala His Leu Phe Val Leu His Pro Tyr Asp Lys Thr Gln Ala
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Tyr Ile Leu Thr Arg Ser Thr Gln His Trp Arg Thr Ser Asn Arg Gly
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Glu Thr Trp Gln Ser Phe Ser Thr Pro His Pro Pro Thr Thr Leu Lys
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Cys Pro Val Ser Ala Pro Pro Ser Ser Pro Pro Glu Gly Lys Thr Trp
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                                 25
Gln Arg His Ser Asp Asn Ala Pro Gln Glu Val Lys Ser Ser Leu Ser
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Asp His Gly Arg Arg Ala Ser Ala Gln Gly Glu Leu Gly Thr Ser Gln
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Ala Thr Pro Pro Arg Ser Met Pro Pro Pro Val Ser Ser Ala Ser Ser
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